# 2025 PRIOR AUTHORIZATION CRITERIA

# **TABLE OF CONTENTS**

Abilify	47
Abilify Mycite Maintenance Kit	47
Abilify Mycite Starter Kit	47
Abiraterone Acetate	491
Abrilada	78
Actemra	79
Actemra Actpen	79
Acthar	
Actimmune	22
Acyclovir	23
Adalimumab-Aacf (2 Pen)	
Adalimumab-Aacf (2 Syringe)	101
Adalimumab-Aacf Starter Pack/Cd/Uc/Hs (6 Pen)	
Adalimumab-Aacf Starter Pack/Psoriasis/Uveitis (4 Pen)	
Adalimumab-Aaty 1-Pen Kit	
Adalimumab-Aaty 2-Pen Kit	
Adalimumab-Aaty 2-Syringe Kit	
Adalimumab-Adaz	
Adalimumab-Fkjp	95
Adapalene	
Adapalene Pump	
Adbry	
Adcirca	453
Adempas	440
Afinitor	
Afinitor Disperz	491
Agamree	
Aimovig	
Ajovy	27
Akeega	
Aklief	
Alcohol Swabs	28
Alecensa	491
Alosetron Hcl	29
Altreno	542
Alunbrig	491
Alvaiz	33
Alyg	
Ambrisentan	
Amjevita	
Amnyra	

Ancobon	241
Androgel Pump	45
Apokyn	49
Apomorphine Hcl	49
Aptensio Xr	50
Aralast Np	30
Aranesp Albumin Free	
Arazlo	541
Arcalyst	
Arikayce	
Aripiprazole	
Aripiprazole Odt	
Armodafinil	
Asceniv	
Ascomp/Codeine	
Asenapine Maleate Sl	
Ativan	
Atovaquone	
Atralin	
Aubagio	
Augtyro	
Austedo	
Austedo Xr	
Austedo XI	
Aveed	
Avita	
Avonex	
Avonex Pen	
Ayvakit	
Azstarys	
Bafiertam	
Balversa	
Belbuca	
Belsomra	
Benlysta SC	
Benztropine Mesylate	
Berinert	
Besremi	
Betaseron	
Bethkis	
Bexarotene Capsule	
Bexarotene Gel	
Bimzelx	82
Bivigam	160
Bonjesta	297
Bosentan	444
Bosulif	491
Braftovi	491

Brukinsa	491
Budesonide Dr (Entocort)	161
Budesonide Er (Uceris)	162
Buphenyl	557
Buprenorphine	
Butalbital/Acetaminophen/Caffeine/Codeine	
Butalbital/Aspirin/Caffeine/Codeine	
Butrans	
Bydureon Bcise	
Byetta	
Bylvay	
Bylvay (Pellets)	
Cabometyx	
Calquence	
Camzyos	
•	
Caprelsa	
Carbaglu	
Carbinoxamine Maleate	
Carglumic Acid	
Cayston	
Cequa	
Cerdelga	
Cerezyme	
Chenodal	169
Chlordiazepoxide Hcl	65
Chlordiazepoxide/Amitriptyline	66
Chlorpromazine Hcl	47
Chorionic Gonadotropin	170
Cialis	63
Cibingo	171
Cimzia	84
Cimzia Starter Kit	84
Cinacalcet Hcl	172
Cingair	
Cinryze	
Clemastine Fumarate	
Clobazam	
Clorazepate Dipotassium	
Clozapine	
Clozapine Odt	
Clozaril	
Cometriq	
Concerta	
Conzip	
Considera	
Copiktra	
Corlanor	
Cortrophin	189

Cosentyx	86
Cosentyx Sensoready Pen	86
Cosentyx Unoready	86
Cotellic	491
Cotempla Xr-Odt	191
Cresemba	192
Crinone	193
Crysvita	194
Cutaquig	196
Cuvrior	197
Cyltezo	88
Cyltezo Starter Package For Crohns Disease/Uc/Hs	88
Cyltezo Starter Package For Psoriasis	
Cyltezo Starter Package For Psoriasis/Uveitis	88
Cyproheptadine Hcl	297
Cystadrops	198
Cystagon	200
Cystaran	199
Dalfampridine Er	202
Daliresp	487
Danazol	37
Daraprim	462
Daurismo	491
Daybue	203
Daytrana	345
Dayvigo	204
Deferasirox (Exjade)	313
Deferasirox (Jadenu)	314
Deferiprone	205
Deflazacort	213
Depo-Testosterone	39
Dexmethylphenidate Hcl (Focalin)	242
Dexmethylphenidate Hcl Er (Focalin Xr)	243
Diazepam	69
Diazepam Intensol	69
Dichlorphenamide	206
Diclegis	297
Diclofenac Epolamine	537
Diclofenac Sodium (Pennsaid)	539
Diclofenac Sodium Gel 3%	535
Dicyclomine Hcl	297
Differin	540
Dihydroergotamine Mesylate Spray	348
Dimethyl Fumarate	
Dimethyl Fumarate Starterpack	
Diphenoxylate/Atropine	
Doptelet	207
Doxepin Hcl	536

Doxylamine Succinate/Pyridoxine Hcl	297
Droxidopa	209
Dupixent	210
Edaravone	470
Elelyso	251
Elidel	57
Eligard	324
Elmiron	212
Emflaza	213
Emgality	214
Emsam	215
Enbrel	90
Enbrel Mini	90
Enbrel Sureclick	90
Endari	217
Enspryng	218
Entyvio Pen	
Eohilia	
Epclusa	
Epidiolex	
Epogen	
Erivedge	
Erleada	
Erlotinib Hcl	
Esbriet	
Estazolam	
Eucrisa	
Evenity	
Everolimus	
Evkeeza	
Evrysdi	
Exjade	
Exkivity	
Extavia	
Eysuvis	
Fabhalta	
Fabior	
Fanapt	
Fanapt Titration Pack	
Fasenra	
Fasenra Pen	
Fentanyl	
Fentanyl Citrate Oral Transmucosal	
Fentanyl Citrate Oral Transmucosal	
Fentora	
Ferriprox	
Ferriprox Twice-A-Day	
Filsnari	237

Filsuvez	238
Fingolimod	355
Fintepla	239
Fioricet/Codeine	297
Firazyr	285
Firdapse	
Flebogamma Dif	
Flector	
Flucytosine	
Fluphenazine Decanoate	
Fluphenazine Hcl	
Flurazepam Hcl	
Focalin	
Focalin Xr	
Forteo	
Fotivda	
Fruzagla	
Fulphila	
Fylnetra	
•	
Gabapentin Once-DailyGalafold	
Gamastan	
Gammagard Liquid	
Gammagard S/D Iga Less Than 1Mcg/MI	
Gammaked	
Gammaplex	
Gamunex-C	
Gattex	
Gauze Pads	
Gavreto	
Gefitinib	
Genotropin	
Genotropin Miniquick	
Geodon	
Gilenya	
Gilotrif	
Glassia	
Glatiramer Acetate	
Glatopa	
Gleevec	
Gocovri	35
Gralise	256
Granix	176
Grastek	411
Hadlima	93
Hadlima Pushtouch	93
Haegarda	283
Haldol Decanoate 100	47

Haldol Decanoate 50	47
Haloperidol	47
Haloperidol Decanoate	47
Haloperidol Lactate	47
Harvoni	295
Hetlioz	524
Hetlioz Lq	296
Horizant	301
Hulio	95
Humatrope	260
Humira	97
Humira Pen	97
Humira Pen-Cd/Uc/Hs Starter	97
Humira Pen-Pediatric Uc Starter Pack	97
Humira Pen-Ps/Uv Starter	97
Hydrocodone Bitartrate Er	395
Hydromorphone Hcl Er	397
Hydroxyzine Hcl	297
Hydroxyzine Pamoate	297
Hyftor	303
Hyrimoz	99
Hyrimoz Crohns Disease And Ulcerative Colitis Starter Pack	99
Hyrimoz Pediatric Crohns Disease Starter Pack	99
Hyrimoz Plaque Psoriasis/Uveitis Starter Pack	99
Hyrimoz Sensoready Pen	99
Hysingla Er	395
Ibrance	491
Icatibant Acetate	285
Iclusig	491
Idacio (2 Pen)	101
Idacio (2 Syringe)	101
Idacio Starter Package For Crohns Disease	101
Idacio Starter Package For Plaque Psoriasis	101
Idhifa	491
llumya	
Imatinib Mesylate	
Imbruvica	
Imiquimod	305
Imiquimod (Zyclara)	598
Imiguimod Pump	
Inbrija	
Inflectra	
Ingrezza	
Inlyta	
Ingovi	
Inrebic	
Insulin Pen Needle	
Insulin Syringe/Needle	

Intrarosa	312
Invega	47
Iressa	491
Isturisa	315
Ivabradine	188
Ivermectin Cream	316
Ivermectin Tablet	317
lwilfin	
Jadenu	
Jadenu Sprinkle	
Jakafi	
Jatenzo	
Javygtor	
Jaypirca	
Joenja	
Jornay Pm	
Juxtapid	
Jynarque	
Kalbitor	
Kalydeco	
•	
Kanjinti	
Kerendia	
Kesimpta	
Keveyis	
Kevzara	
Kineret	
Kisqali	
Kisqali Femara 200 Dose	492
Kisqali Femara 400 Dose	
Kisqali Femara 600 Dose	492
Kitabis Pak	534
Korlym	347
Koselugo	492
Krazati	492
Kuvan	490
Lapatinib Ditosylate	
Latuda	
Lazcluze	
Ledipasvir/Sofosbuvir	
Lenalidomide	
Lenvima 10 Mg Daily Dose	
Lenvima 12Mg Daily Dose	
Lenvima 14 Mg Daily Dose	
Lenvima 18 Mg Daily Dose	
<b>G</b> ,	
Lenvima 20 Mg Daily Dose	
Lenvima 24 Mg Daily Dose	
Lenvima 4 Mg Daily Dose	
Lenvima 8 Mg Daily Dose	492

Letairis	442
Leukine	177
Leuprolide Acetate	324
L-Glutamine	217
Licart	
Lidocaine Ointment	
Lidocaine Patch	
Lidocaine Solution	
Lidocaine/Prilocaine	
Lidocan	
Lidoderm	
Linezolid	
Ligrev	
·	
Liraglutide	
Litfulo	
Livmarli	
Lomotil	
Lonsurf	
Lorazepam	
Lorazepam Intensol	
Lorbrena	492
Loreev Xr	72
Lotronex	29
Loxapine	47
Lumakras	492
Lumryz	334
Lupkynis	335
Lupron Depot (1-Month)	324
Lupron Depot (3-Month)	
Lupron Depot (4-Month)	
Lupron Depot (6-Month)	
Lupron Depot-Ped (1-Month)	
Lupron Depot-Ped (3-Month)	
Lupron Depot-Ped (6-Month)	
Lurasidone Hcl	
Lybalvi	
,	
Lynparza	
Lytgobi	
Matulane	
Mavenclad	
Mavyret	
Mayzent	
Mayzent Starter Pack	
Mekinist	
Mektovi	
Memantine Hcl	338
Memantine Hcl Er	337
Memantine Hcl Titration Pak	338

Mepron	59
Metadate Cd	341
Methamphetamine Hcl	339
Methitest	44
Methylin	340
Methylphenidate Hcl (Methylin)	340
Methylphenidate Hcl (Ritalin)	485
Methylphenidate Hcl Cd	341
Methylphenidate Hcl Chewable	342
Methylphenidate Hcl Er (Aptensio Xr)	50
Methylphenidate Hcl Er (Concerta)	187
Methylphenidate Hcl Er (La)	344
Methylphenidate Hcl Er (Relexxii)	
Methylphenidate Hcl Er (Ritalin LA)	
Methylphenidate Hcl Er Capsule	
Methylphenidate Hcl Er Tablet	
Methylphenidate Patch	
Methyltestosterone	
Miebo	
Mifepristone	
Miglustat	
Migranal	
Modafinil	
Molindone Hcl	
Morphine Sulfate Er	
Mounjaro	
Ms Contin	
Mulpleta	
Mvasi	
Myalept	
Mycapssa	
Myfembree	
Namenda	
Namenda Titration Pak	
Namenda Xr	
Natesto	
Nerlynx	
Neulasta	
Neulasta Onpro Kit	
Neupogen	
Nexavar	
Nexietol	
Nexizet	
Nexilzet	
Ninlaro	
Nillaro	
Nocdurna	
Norditropin Flexpro	
NOTOTO CONT. FIEXULO	

Northera	209
Nourianz	376
Novarel	170
Noxafil	432
Nubega	492
Nucala	
Nucynta Er	
Nuedexta	
Nuplazid	
Nurtec	
Nutropin Aq Nuspin 10	
Nutropin Aq Nuspin 20	
Nutropin Aq Nuspin 5	
Nuvigil	
Nyvepria	
Ocaliva	
Octagam	
Octreotide Acetate	
Odactra	
Odomzo	
Ofev	
Ogsiveo	
Ojemda	
Ojjaara	
Olanzapine	
Olanzapine Odt	
Olanzapine/Fluoxetine	
Olpruva	
Olumiant	
Omnipod 5 Kit	
Omnipod 5 Pods	386
Omnipod Classic Kit	386
Omnipod Classic Pods	386
Omnipod Dash Kit	386
Omnipod Dash Pods	386
Omnipod Go	386
Omnitrope	270
Omvoh	114
Onfi	67
Ontruzant	308
Onureg	492
Opsumit	
Opsynvi	
Opzelura	
Oralair	
Orencia	
Orencia Clickject	
Orenitram	

Orenitram Titration Kit Month 1	449
Orenitram Titration Kit Month 2	449
Orenitram Titration Kit Month 3	449
Orgovyx	492
Oriahnn	
Orilissa	415
Orkambi	
Orladeyo	
Ormalvi	
Orserdu	
Osmolex Er	
Osphena	
Otezla	
Otezia Otrexup	
OttexupOttexup	
Oxbryta	
•	
Oxervate	
Oxlumo	
Oxycodone Hcl Er	
Oxycontin	
Oxymorphone Hcl Er	
Ozempic	
Palforzia	
Paliperidone Er	47
Palynziq	427
Panretin	428
Panzyga	429
Pazopanib Hcl	492
Pegasys	430
Pemazyre	492
Pennsaid	539
Perphenazine	47
· Pheburane	
Pimecrolimus	
Pimozide	
Pigray 200Mg Daily Dose	
Pigray 250Mg Daily Dose	
Pigray 300Mg Daily Dose	
Pirfenidone	
Plegridy	
g ,	
Plegridy Starter Pack	
Pliaglis	
Pomalyst	
Ponvory	
Ponvory 14-Day Starter Pack	
Posaconazole Dr	
Posaconazole Inj	
Posaconazole Susp	432

Praluent	434
Pregnyl	170
Privigen	160
Procrit	
Procysbi	201
Prolastin-C	
Prolia	435
Promacta	438
Promethazine Hcl	297
Promethazine Hcl Plain	297
Promethazine Vc	297
Promethazine/Phenylephrine	297
Promethegan	297
Provigil	
Prudoxin	536
Pyrimethamine	462
, Pyrukynd	463
Pyrukynd Taper Pack	
, , , , , , , , , , , , , , , , , , ,	
Qinlock	
Qualaquin	
Quetiapine Fumarate	
Quetiapine Fumarate Er	
Quillichew Er	
Quillivant Xr	
Quinine Sulfate	
Qulipta	
Quvivig	
Radicava	
Radicava Ors	
Radicava Ors Starter Kit	
Rasuvo	
Ravicti	
Rayos	
Rebif	
Rebif Rebidose	
Rebif Rebidose Titration Pack	
Rebif Titration Pack	
Reblozyl	
Recorlev	
Regranex	
Releuko	
Relexxii	
Relistor Injection	
Relistor Tablet	
Remicade	
Renflexis	
Repatha	
NCPULIU	<del>4</del> 0U

Repatha Pushtronex System	480
Repatha Sureclick	480
Retacrit	225
Retevmo	493
Retin-A	542
Retin-A Micro	542
Retin-A Micro Pump	
Revatio	
Revlimid	
Rexulti	
Reyvow	
Rezdiffra	
Rezlidhia	
Rezurock	
Riabni	
Rinvoq Lq	
Rinvoq Tablet	
Risperdal	
Risperidone Odt	
Ritalin	
Ritalin La	
Rivfloza	
Roflumilast	
Rozlytrek	
Rubraca	
Ruconest	
Ruxience	
Rybelsus	
Rydapt	
Ryvent	
Sajazir	
Samsca	489
Sandostatin	506
Sandostatin Lar Depot	508
Saphris	47
Sapropterin Dihydrochloride	490
Scemblix	493
Scopolamine	297
Secuado	47
Sensipar	172
Seroquel	47
Seroquel Xr	
Serostim	
Signifor	
Signifor Lar	
Sildenafil Citrate	
Silig	
Simlandi 1Pn Kit	

Simlandi 2Pn Inj	132
Simponi	133
Sivextro	499
Skyclarys	501
Skyrizi	135
Skyrizi Pen	
, Skytrofa	
Sodium Oxybate	
Sodium Phenylbutyrate	
Sofosbuvir/Velpatasvir	
Sogroya	
Somatuline Depot	
Somavert	
Soolantra	
Sorafenib	
Sotyktu	
Sovaldi	
Spevigo	
Sprycel	
Stelara	
Stimufend	
Stivarga	
Strensiq	
Stromectol	
Sucraid	
Sunitinib Malate	
Sunosi	
Sutent	
Symbyax	
Symdeko	
Sympazan	
Symproic	
Syprine	
T Tretinoin Cream, Gel	
Tabrecta	
Tacrolimus	
Tadalafil 2.5Mg, 5Mg	63
Tadalafil Tablet 20Mg	453
Tadliq	453
Tafinlar	493
Tagrisso	493
Takhzyro	
Taltz	141
Talzenna	493
Targretin Capsule	493
Targretin Gel	76
Tarpeyo	523
Tascenso Odt	363

Tasigna	493
Tasimelteon	524
Tavalisse	525
Tavneos	526
Tazarotene	541
Tazorac	541
Tazverik	493
Tecfidera	354
Tecfidera Starter Pack	354
Tegsedi	527
Tepmetko	493
Teriflunomide	
Teriparatide	
Testim	
Testosterone	
Testosterone Cypionate	
Testosterone Enanthate	
Testosterone Pump	
Testosterone Topical Solution	
Tetrabenazine	
Tezspire	
Thalomid	
Thioridazine Hcl	
Thiothixene	
Tibsovo	
Tlando	
Tobi	
Tobi Podhaler	
Tobramycin (Bethkis)	
Tobramycin Neb	
Tolvaptan	
Torpenz	
Tracleer	
Tramadol Hcl Er	
Transderm-Scop	
Trazimera	
Trelstar Mixject	
Tremfya	
Tretinoin Capsule 10Mg	
Tretinoin Microsphere	
Tretinoin Microsphere Pump	
Tridacaine II	
Trientine Hcl	
Trifluoperazine Hcl	
Trihexyphenidyl Hcl	
Trikafta	
Trudhesa	
Trulicity	548

Truqap	493
Tukysa	493
Turalio	493
Tyenne Inj	145
Tykerb	493
, Tymlos	550
, Tyrvaya	
Tyvaso DPI Institutional Kit	
Tyvaso DPI Maintenance Kit	
Tyvaso DPI Titration Kit	
Ubrelvy	
Uceris	
Udenyca	
Udenyca Onbody	
Uptravi	
Uptravi Titration Pack	
Valchlor	
Valium	
Vanflyta	
Vannyta Velsipity	
• •	
Venclexta	
Venclexta Starting Pack	
Ventavis	
Veozah	
Verkazia	
Versacloz	
Verzenio	493
Vevye	389
Vfend	567
Vfend lv	567
V-Go	561
Viberzi	562
Victoza	563
Vijoice	565
Vistaril	
Vitrakvi	
Vivjoa	
Vizimpro	
Vogelxo	
Vogelxo Pump	
Vonjo	
Voranigo	
Voriconazole	
Vosevi	
Votrient	
Vowst	
Voxzogo	
Vovdeva	572

Vpriv	253
Vtama	573
Vumerity	365
Vyndamax	574
Vyndaqel	575
Wainua	576
Wakix	577
Wegovy	578
Welireg	493
Winlevi	579
Winrevair	461
Xalkori	493
Xdemvy	580
Xeljanz Solution	
Xeljanz Tablet	
Xeljanz Xr	
Xembify	
Xenazine	
Xermelo	
Xgeva	
Xifaxan	
Xiidra	
Xolair	
Xolremdi	
Xospata	
Xpovio	
Xtampza Er	
Xtandi	
Xyosted	
Xyrem	
Xywav	
Yargesa	
Yonsa	
Yuflyma 1-Pen Kit	_
•	
Yuflyma 2-Pen Kit	
Yuflyma 2-Syringe Kit	
Yuflyma Cd/Uc/Hs Starter	
Yusimry	
Zarxio	
Zavesca	
Zavzpret	
Zejula	
Zelboraf	
Zemaira	
Zepatier	
Zeposia	
Zeposia 7-Day Starter Pack	
Zeposia Starter Kit	366

Ziextenzo	186
Zilbrysq	593
Ziprasidone Mesylate	48
Zirabev	308
Zokinvy	594
Zolinza	494
Zomacton	276
Zonalon	536
Zoryve Cream	595
Zoryve Foam	596
Zovirax	23
Ztalmy	597
Ztlido	330
Zyclara	598
Zyclara Pump	598
Zydelig	494
Zykadia	494
Zymfentra 1-Pen	158
Zymfentra 2-Pen	158
Zymfentra 2-Syringe	158
Zyprexa	48
Zyprexa Relprevv	48
Zyprexa Zydis	48
	494
, c Zvvox	

Acthar Gel PA

### Drug Name(s)

Acthar

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of infantile spasm OR
  - B. Patient has a diagnosis of nephrotic syndrome AND ONE of the following:
    - i. Patient has failed a conventional agent (i.e., prednisone, tacrolimus) for the requested indication OR
    - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to a conventional agent OR
  - C. Patient has a diagnosis of multiple sclerosis AND ALL of the following:
    - i. Patient is experiencing an acute exacerbation AND
    - ii. If indicated, there is evidence of a claim that the patient is currently being treated with a disease modifying drug (DMD) within the past 90 days [e.g., Avonex, dimethyl fumarate, glatiramer] to control disease progression OR has an intolerance, FDA labeled contraindication, or hypersensitivity to a DMD AND
    - iii. ONE of the following:
      - 1. Patient has failed corticosteroid therapy (e.g., methylprednisolone) within the last 30 days OR
      - 2. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to corticosteroid therapy OR

Criteria continues: see Other Criteria

### **Age Restriction:**

For diagnosis of infantile spasm, patient is less than 2 years of age.

# **Prescriber Restrictions:**

# **Coverage Duration:**

6 months for infantile spasm, 1 month for all other indications

- D. Patient has a diagnosis of rheumatic disorder (e.g., ankylosing spondylitis, juvenile idiopathic arthritis, juvenile rheumatoid arthritis, psoriatic arthritis, rheumatoid arthritis) AND ALL of the following:
  - i. The requested agent will be used as adjunct therapy for short-term administration (to tide the patient over an acute episode or exacerbation) AND

- ii. There is evidence of a claim that the patient is currently being treated with a conventional agent within the past 90 days [e.g., DMARD (methotrexate, leflunomide), biologics (Hadlima)] to control disease progression AND
- iii. ONE of the following:
  - 1. Patient has failed corticosteroid therapy (e.g., methylprednisolone) within the last 30 days

OR

- 2. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to corticosteroid therapy OR
- E. Patient has a diagnosis of systemic lupus erythematosus (SLE) disease AND the patient will continue standard SLE therapy [corticosteroids (e.g., methylprednisolone, prednisone), hydroxychloroquine, immunosuppressives (e.g., azathioprine, methotrexate, oral cyclophosphamide)] in combination with the requested agent OR
- F. Patient has another FDA approved indication AND ONE of the following:
  - i. Patient has failed corticosteroid therapy (e.g., methylprednisolone) within the last 30 days OR
  - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to corticosteroid therapy OR
- G. Patient has another indication that is supported in CMS approved compendia for the requested agent AND ONE of the following:
  - i. Patient has failed corticosteroid therapy (e.g., methylprednisolone) within the last 30 days OR
  - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to corticosteroid therapy AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Actimmune PA

Drug Name(s)

Actimmune

**Indications:** 

All Medically-Accepted Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Acyclovir Topical PA

Drug Name(s)

Acyclovir

Zovirax

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Adbry PA

### Drug Name(s)

Adbry

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of moderate-to-severe atopic dermatitis AND BOTH of the following:
  - A. ONE of the following:
    - i. Patient has tried and failed a topical steroid (e.g., triamcinolone) OR
    - ii. Patient has an intolerance or hypersensitivity to a topical steroid OR
    - iii. Patient has an FDA labeled contraindication to a topical steroid AND
  - B. ONE of the following:
    - i. Patient has tried and failed a topical calcineurin inhibitor (e.g., tacrolimus) OR
    - ii. Patient has an intolerance or hypersensitivity to a topical calcineurin inhibitor OR
    - iii. Patient has an FDA labeled contraindication to a topical calcineurin inhibitor AND
- 2. Patient will NOT be using the requested agent in combination with another biologic agent or a JAK inhibitor for the requested indication AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of moderate-to-severe atopic dermatitis AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another biologic agent or a JAK inhibitor for the requested indication AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

### **Age Restriction:**

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, dermatologist, immunologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Agamree PA

### Drug Name(s)

Agamree

#### Indications:

All FDA-Approved Indications.

#### **Off-Label Uses:**

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require the following:

- 1. Patient has a diagnosis of Duchenne muscular dystrophy (DMD) confirmed by ONE of the following:
  - A. Presence of abnormal dystrophin OR
  - B. Confirmed mutation of the dystrophin gene

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Duchenne muscular dystrophy (DMD) AND
- 3. Patient has had improvement, stabilization of the disease, or clinical benefit from baseline (e.g., slowed disease progression, improved strength and timed motor function, improved pulmonary function, reduced the need for scoliosis surgery)

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be for 12 months

Aimovig PA

### Drug Name(s)

Aimovig

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of migraine AND
- 2. The requested agent is being used for migraine prophylaxis AND
- 3. Patient has 4 or more migraine headache days per month AND
- 4. Patient will NOT be using the requested agent in combination with another calcitonin gene-related peptide (CGRP) agent for migraine prophylaxis

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of migraine AND
- 3. The requested agent is being used for migraine prophylaxis AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with another calcitonin gene-related peptide (CGRP) agent for migraine prophylaxis

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Ajovy PA

### Drug Name(s)

Ajovy

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of migraine AND
- 2. The requested agent is being used for migraine prophylaxis AND
- 3. Patient has 4 or more migraine headache days per month AND
- 4. Patient will NOT be using the requested agent in combination with another calcitonin gene-related peptide (CGRP) agent for migraine prophylaxis

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of migraine AND
- 3. The requested agent is being used for migraine prophylaxis AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with another calcitonin gene-related peptide (CGRP) agent for migraine prophylaxis

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Alcohol Swabs PA

# Drug Name(s)

Alcohol Swabs

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

This program will be implemented as a dynamic PA.

Criteria for approval require BOTH of the following:

- 1. The requested medical supply product will be used in the delivery of insulin to the body AND
- 2. Patient's medication history includes use of insulin within the past 180 days

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Alosetron PA

### Drug Name(s)

Alosetron Hcl

Lotronex

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of irritable bowel syndrome with severe diarrhea (IBS-D) AND
- 2. Patient's sex is female AND
- 3. Patient exhibits at least ONE of the following:
  - a. Frequent and severe abdominal pain/discomfort OR
  - b. Frequent bowel urgency or fecal incontinence OR
  - c. Disability or restriction of daily activities due to IBS AND
- 4. Prescriber has ruled out anatomic or biochemical abnormalities of the gastrointestinal tract

### Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Alpha-1-Proteinase Inhibitor PA - Aralast/Zemaira

# Drug Name(s)

Aralast Np

Zemaira

#### **Indications:**

All FDA-Approved Indications.

#### **Off-Label Uses:**

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of alpha-1 antitrypsin deficiency (AATD) with clinically evident emphysema AND
- 2. Patient has a pre-treatment serum alpha-1 antitrypsin (AAT) level less than 11 micromol/L (80 mg/dL by immunodiffusion or 57 mg/dL using nephelometry) AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of alpha-1 antitrypsin deficiency (AATD) with clinically evident emphysema AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 months

Alpha-1-Proteinase Inhibitor PA - Glassia

### Drug Name(s)

Glassia

#### Indications:

All FDA-Approved Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of alpha-1 antitrypsin deficiency (AATD) with clinically evident emphysema AND
- 2. Patient has a pre-treatment serum alpha-1 antitrypsin (AAT) level less than 11 micromol/L (80 mg/dL by immunodiffusion or 57 mg/dL using nephelometry) AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of alpha-1 antitrypsin deficiency (AATD) with clinically evident emphysema AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 months

Alpha-1-Proteinase Inhibitor PA - Prolastin-C

### Drug Name(s)

Prolastin-C

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of alpha-1 antitrypsin deficiency (AATD) with clinically evident emphysema AND
- 2. Patient has a pre-treatment serum alpha-1 antitrypsin (AAT) level less than 11 micromol/L (80 mg/dL by immunodiffusion or 57 mg/dL using nephelometry) AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of alpha-1 antitrypsin deficiency (AATD) with clinically evident emphysema AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 months

Alvaiz PA

### Drug Name(s)

Alvaiz

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of persistent or chronic immune (idiopathic) thrombocytopenia (ITP) AND ONE of the following:
    - i. Patient has tried and had an insufficient response to a corticosteroid or immunoglobulin (IVIg or anti-D) OR
    - ii. Patient has an intolerance or hypersensitivity to a corticosteroid or immunoglobulin (IVIg or anti-D) OR
    - iii. Patient has an FDA labeled contraindication to a corticosteroid or immunoglobulin (IVIg or anti-D) OR
    - iv. Patient has had an insufficient response to a splenectomy OR
  - B. Patient has a diagnosis of hepatitis C associated thrombocytopenia AND ONE of the following:
    - i. Patient's platelet count is less than 75 x  $10^9/L$  AND the intent is to increase platelet counts sufficiently to initiate interferon therapy OR
    - ii. Patient is on concomitant therapy with interferon therapy AND is at risk for discontinuing hepatitis C therapy due to thrombocytopenia OR
  - C. Patient has a diagnosis of severe aplastic anemia (SAA) AND ALL of the following:
    - i. Patient has at least 2 of the following blood criteria:
      - 1. Neutrophils less than 0.5 X 10^9/L OR
      - 2. Platelets less than 30 X 10<sup>9</sup>/L OR
      - 3. Reticulocyte count less than 60 X 10^9/L AND
    - ii. Patient has at least 1 of the following marrow criteria:
      - 1. Severe hypocellularity is less than 25% OR
      - 2. Moderate hypocellularity is 25-50% with hematopoietic cells representing less than 30% of residual cells AND
    - iii. Patient has tried and had an insufficient response to BOTH antithymocyte globulin (ATG) AND cyclosporine therapy OR
  - D. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

#### Age Restriction:

### **Prescriber Restrictions:**

# **Coverage Duration:**

Initial: 6 months for ITP. Renewal: 12 months for ITP. Other indications, see Other Criteria.

#### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of persistent or chronic immune (idiopathic) thrombocytopenia (ITP) AND ONE of the following:
    - i. Patient's platelet count is 50 x 10^9/L or greater OR
    - ii. Patient's platelet count has increased sufficiently to avoid clinically significant bleeding OR
  - B. Patient has a diagnosis of hepatitis C associated thrombocytopenia AND BOTH of the following:
    - i. ONE of the following:
      - 1. Patient will be initiating hepatitis C therapy with interferon therapy OR
      - 2. Patient will be maintaining hepatitis C therapy with interferon therapy at the same time as the requested agent AND
    - ii. ONE of the following:
      - 1. Patient's platelet count is 90 x 10^9/L or greater OR
      - 2. Patient's platelet count has increased sufficiently to initiate or maintain interferon therapy for the treatment of hepatitis C OR
  - C. Patient has a diagnosis of severe aplastic anemia (SAA) AND the patient has had clinical benefit with the requested agent OR
  - D. Patient has another indication that is supported in CMS approved compendia AND the patient has had clinical benefit with the requested agent AND
- 3. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Initial: 48 weeks for hepatitis C associated thrombocytopenia, 16 weeks for SAA, 12 months for All other indications

Renewal: 48 weeks for hepatitis C associated thrombocytopenia, 12 months for SAA, 12 months for All other indications

Amantadine ER PA – Gocovri

# Drug Name(s)

Gocovri

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of Parkinson's disease AND
- 2. ONE of the following:
  - A. The requested agent will be used for the treatment of dyskinesia OR
  - B. The requested agent will be used for the adjunctive treatment in patients experiencing "off" episodes AND
- 3. The requested agent will be used in combination with levodopa-based therapy

### **Age Restriction:**

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Amantadine ER PA - Osmolex ER

# Drug Name(s)

Osmolex Er

# **Indications:**

All FDA-Approved Indications.

#### **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of Parkinson's disease OR
  - B. BOTH of the following:
    - i. Patient has a diagnosis of drug-induced extrapyramidal reaction AND
    - ii. Prescriber has assessed and adjusted, if applicable, any medications known to cause extrapyramidal symptoms

### **Age Restriction:**

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Anabolic Steroid PA - Danazol

# Drug Name(s)

Danazol

### **Indications:**

All Medically-Accepted Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. Patient will NOT be using the requested agent in combination with another androgen or anabolic steroid OR
  - B. Prescriber has provided information in support of therapy with more than one agent

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Androgen Injectable PA - Aveed

### Drug Name(s)

Aveed

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient's sex is male with a diagnosis of primary or secondary (hypogonadotropic) hypogonadism AND
- 2. ONE of the following:
  - A. Patient is NOT currently receiving testosterone replacement therapy AND has ONE of the following pretreatment levels:
    - i. Total serum testosterone level that is below the testing laboratory's lower limit of the normal range or is less than 300 ng/dL OR
    - ii. Free serum testosterone level that is below the testing laboratory's lower limit of the normal range OR
  - B. Patient is currently receiving testosterone replacement therapy AND has ONE of the following current levels:
    - i. Total serum testosterone level that is within the testing laboratory's normal range OR below the testing laboratory's lower limit of the normal range OR is less than 300  $\,$  ng/dL  $\,$  OR
    - ii. Free serum testosterone level is within the testing laboratory's normal range OR below the testing laboratory's normal range AND
- 3. ONE of the following:
  - A. Patient will NOT be using the requested agent in combination with another androgen or anabolic steroid OR
  - B. Prescriber has provided information in support of therapy with more than one agent

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Androgen Injectable PA - testosterone cypionate

# Drug Name(s)

Depo-Testosterone

Testosterone Cypionate

#### **Indications:**

All Medically-Accepted Indications.

#### **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Patient's sex is male with AIDS/HIV-associated wasting syndrome AND BOTH of the following:
    - i. ONE of the following:
      - a. Unexplained involuntary weight loss (greater than 10% baseline body weight within 12 months, or 7.5% within 6 months) OR
      - b. Body mass index less than 20 kg/m2 OR
      - c. At least 5% total body cell mass (BCM) loss within 6 months OR
      - d. BCM less than 35% of total body weight and BMI less than 27 kg/m2 AND
    - ii. All other causes of weight loss have been ruled out OR
  - B. Patient's sex is female with metastatic/inoperable breast cancer OR
  - C. Patient's sex is male with primary or secondary (hypogonadotropic) hypogonadism OR
  - D. Patient's sex is male and is an adolescent with delayed puberty AND
- 2. If the patient's sex is a male, ONE of the following:
  - A. Patient is NOT currently receiving testosterone replacement therapy AND has ONE of the following pretreatment levels:
    - i. Total serum testosterone level that is below the testing laboratory's lower limit of the normal range or is less than 300 ng/dL OR
    - ii. Free serum testosterone level that is below the testing laboratory's lower limit of the normal range OR
  - B. Patient is currently receiving testosterone replacement therapy AND has ONE of the following current levels:
    - i. Total serum testosterone level that is within the testing laboratory's normal range OR below the testing laboratory's lower limit of the normal range OR is less than 300 ng/dL OR  $\,$
    - ii. Free serum testosterone level is within the testing laboratory's normal range OR below the testing laboratory's normal range AND
- 3. ONE of the following:
  - A. Patient will NOT be using the requested agent in combination with another androgen or anabolic steroid OR
  - B. Prescriber has provided information in support of therapy with more than one agent

# **Age Restriction:**

### **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be 6 months for delayed puberty, 12 months for all other indications **Other Criteria:** 

Androgen Injectable PA - testosterone enanthate

# Drug Name(s)

Testosterone Enanthate

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Patient's sex is male with AIDS/HIV-associated wasting syndrome AND BOTH of the following:
    - i. ONE of the following:
      - a. Unexplained involuntary weight loss (greater than 10% baseline body weight within 12 months, or 7.5% within 6 months) OR
      - b. Body mass index less than 20 kg/m2 OR
      - c. At least 5% total body cell mass (BCM) loss within 6 months OR
      - d. BCM less than 35% of total body weight and BMI less than 27 kg/m2 AND
    - ii. All other causes of weight loss have been ruled out OR
  - B. Patient's sex is female with metastatic/inoperable breast cancer OR
  - C. Patient's sex is male with primary or secondary (hypogonadotropic) hypogonadism OR
  - D. Patient's sex is male and is an adolescent with delayed puberty AND
- 2. If the patient's sex is a male, ONE of the following:
  - A. Patient is NOT currently receiving testosterone replacement therapy AND has ONE of the following pretreatment levels:
    - i. Total serum testosterone level that is below the testing laboratory's lower limit of the normal range or is less than 300 ng/dL OR
    - ii. Free serum testosterone level that is below the testing laboratory's lower limit of the normal range OR
  - B. Patient is currently receiving testosterone replacement therapy AND has ONE of the following current levels:
    - i. Total serum testosterone level that is within the testing laboratory's normal range OR below the testing laboratory's lower limit of the normal range OR is less than 300 ng/dL OR
    - ii. Free serum testosterone level is within the testing laboratory's normal range OR below the testing laboratory's normal range AND
- 3. ONE of the following:
  - A. Patient will NOT be using the requested agent in combination with another androgen or anabolic steroid OR
  - B. Prescriber has provided information in support of therapy with more than one agent

# **Age Restriction:**

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be 6 months for delayed puberty, 12 months for all other indications  Other Criteria:

Androgen Injectable PA - Xyosted

### Drug Name(s)

**Xyosted** 

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Patient's sex is male with AIDS/HIV-associated wasting syndrome AND BOTH of the following:
    - i. ONE of the following:
      - a. Unexplained involuntary weight loss (greater than 10% baseline body weight within 12 months, or 7.5% within 6 months) OR
      - b. Body mass index less than 20 kg/m2 OR
      - c. At least 5% total body cell mass (BCM) loss within 6 months OR
      - d. BCM less than 35% of total body weight and BMI less than 27 kg/m2 AND
    - ii. All other causes of weight loss have been ruled out OR
  - B. Patient's sex is male with primary or secondary (hypogonadotropic) hypogonadism AND
- 2. Patient's sex is male with ONE of the following:
  - A. Patient is NOT currently receiving testosterone replacement therapy AND has ONE of the following pretreatment levels:
    - i. Total serum testosterone level that is below the testing laboratory's lower limit of the normal range or is less than 300 ng/dL OR
    - ii. Free serum testosterone level that is below the testing laboratory's lower limit of the normal range OR
  - B. Patient is currently receiving testosterone replacement therapy AND has ONE of the following current levels:
    - i. Total serum testosterone level that is within the testing laboratory's normal range OR below the testing laboratory's lower limit of the normal range OR is less than 300 ng/dL OR  $\,$
    - ii. Free serum testosterone level is within the testing laboratory's normal range OR below the testing laboratory's normal range AND
- 3. ONE of the following:
  - A. Patient will NOT be using the requested agent in combination with another androgen or anabolic steroid OR
  - B. Prescriber has provided information in support of therapy with more than one agent

#### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Androgen Oral PA

# Drug Name(s)

Methitest

Methyltestosterone

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Patient's sex is male with cryptorchidism OR
  - B. Patient's sex is male with hypogonadism OR
  - C. Patient's sex is male and is an adolescent with delayed puberty OR
  - D. Patient's sex is female with metastatic/inoperable breast cancer AND
- 2. If the patient's sex is male, ONE of the following:
  - A. Patient is NOT currently receiving testosterone replacement therapy AND has ONE of the following pretreatment levels:
    - i. Total serum testosterone level that is below the testing laboratory's lower limit of the normal range or is less than 300 ng/dL OR
    - ii. Free serum testosterone level that is below the testing laboratory's lower limit of the normal range OR
  - B. Patient is currently receiving testosterone replacement therapy AND has ONE of the following current levels:
    - i. Total serum testosterone level that is within the testing laboratory's normal range OR below the testing laboratory's lower limit of the normal range OR is less than 300 ng/dL OR  $\,$
    - ii. Free serum testosterone level is within the testing laboratory's normal range OR below the testing laboratory's normal range AND
- 3. ONE of the following:
  - A. Patient will NOT be using the requested agent in combination with another androgen or anabolic steroid OR
  - B. Prescriber has provided information in support of therapy with more than one agent

# Age Restriction:

#### **Prescriber Restrictions:**

### **Coverage Duration:**

Approval will be 6 months for delayed puberty, 12 months for all other indications

Androgen Topical PA

### Drug Name(s)

**Androgel Pump** 

Natesto

**Testim** 

Testosterone

Testosterone Pump

**Testosterone Topical Solution** 

Vogelxo

Vogelxo Pump

### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Patient has AIDS/HIV-associated wasting syndrome AND BOTH of the following:
    - i. ONE of the following:
      - a. Unexplained involuntary weight loss (greater than 10% baseline body weight within 12 months, or 7.5% within 6 months)  $\rm OR$
      - b. Body mass index less than 20 kg/m2 OR
      - c. At least 5% total body cell mass (BCM) loss within 6 months OR
      - d. In men: BCM less than 35% of total body weight and BMI less than 27 kg/m2 OR
      - e. In women: BCM less than 23% of total body weight and BMI less than 27 kg/m2 AND
    - ii. All other causes of weight loss have been ruled out OR
- B. Patient's sex is male with primary or secondary (hypogonadotropic) hypogonadism AND
- 2. If the patient's sex is male, ONE of the following:
  - A. Patient is NOT currently receiving testosterone replacement therapy AND has ONE of the following pretreatment levels:
    - i. Total serum testosterone level that is below the testing laboratory's lower limit of the normal range or is less than 300 ng/dL OR
    - ii. Free serum testosterone level that is below the testing laboratory's lower limit of the normal range OR
  - B. Patient is currently receiving testosterone replacement therapy AND has ONE of the following current levels:
    - i. Total serum testosterone level that is within the testing laboratory's normal range OR below the testing laboratory's lower limit of the normal range OR is less than 300 ng/dL OR  $\,$

- ii. Free serum testosterone level is within the testing laboratory's normal range OR below the testing laboratory's normal range AND
- 3. ONE of the following:
  - A. Patient will NOT be using the requested agent in combination with another androgen or anabolic steroid OR
  - B. Prescriber has submitted information in support of therapy with more than one agent

# Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Antipsychotics PA

# Drug Name(s)

Abilify

Abilify Mycite Maintenance Kit

Abilify Mycite Starter Kit

Aripiprazole

Aripiprazole Odt

Asenapine Maleate SI

Chlorpromazine Hcl

Clozapine

Clozapine Odt

Clozaril

Fanapt

Fanapt Titration Pack

Fluphenazine Decanoate

Fluphenazine Hcl

Geodon

Haldol Decanoate 100

Haldol Decanoate 50

Haloperidol

Haloperidol Decanoate

Haloperidol Lactate

Invega

Latuda

Loxapine

Lurasidone Hcl

Lvbalvi

Molindone Hcl

Olanzapine

Olanzapine Odt

Olanzapine/Fluoxetine

Paliperidone Er

Perphenazine

Pimozide

Quetiapine Fumarate

Quetiapine Fumarate Er

Rexulti

Risperdal

Risperidone Odt

Saphris

Secuado

Seroquel

Seroquel Xr

Symbyax

Thioridazine Hcl

Thiothixene

Trifluoperazine Hcl

Versacloz

Ziprasidone Mesylate

Zyprexa

Zyprexa Relprevv

Zyprexa Zydis

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

### **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - b. Prescriber states the patient is currently being treated with the requested agent OR
  - c. ONE of the following:
    - i. Patient has a diagnosis other than dementia-related psychosis or dementia related behavioral symptoms OR
    - ii. Patient has dementia-related psychosis or dementia related behavioral symptoms AND BOTH of the following:
      - 1. Dementia-related psychosis is determined to be severe or the associated behavior puts the patient or others in danger AND
      - 2. Prescriber has documented that s/he has discussed the risk of increased mortality with the patient and/or the patient's surrogate decision maker

Approval authorizations will apply to the requested medication only.

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Apomorphine Inj PA

# Drug Name(s)

Apokyn

Apomorphine Hcl

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. The requested agent will be used to treat acute, intermittent hypomobility, "off" episodes ("end of dose wearing off" and unpredictable "on/off" episodes) associated with advanced Parkinson's disease AND
- 2. The requested agent will be used in combination with agents used for therapy in Parkinson's disease (e.g., levodopa, dopamine agonist, monoamine oxidase B inhibitor) AND
- 3. Patient will NOT be using the requested agent in combination with a 5-HT3 antagonist (e.g., ondansetron, granisetron, dolasetron, palonosetron, alosetron)

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Aptensio XR PA

Drug Name(s)

Aptensio Xr

Methylphenidate Hcl Er (Aptensio Xr)

Indications:

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Arcalyst PA

# Drug Name(s)

Arcalyst

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has been diagnosed with Cryopyrin-Associated Periodic Syndromes (CAPS) including Familial Cold Auto-inflammatory Syndrome (FCAS) or Muckle-Wells Syndrome (MWS) OR
  - B. BOTH of the following:
    - i. Patient has a diagnosis of deficiency of interleukin-1 receptor antagonist AND
    - ii. The requested agent is being used for maintenance of remission OR
  - C. BOTH of the following:
    - i. Patient has a diagnosis of recurrent pericarditis AND
    - ii. The requested agent is being used to reduce the risk of recurrence AND
- 2. Patient will NOT be using the requested agent in combination with another biologic agent

### **Age Restriction:**

For diagnosis of CAPS including FCAS or MWS, patient is 12 years of age or over

For diagnosis of recurrent pericarditis and reduction in risk of recurrence, patient is 12 years of age or over

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Arikayce PA

Drug Name(s)

Arikayce

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of Mycobacterium avium complex (MAC) lung disease AND
- 2. Patient has not achieved negative sputum cultures despite at least 6 consecutive months of treatment with standard combination antibiotic therapy for MAC lung disease [e.g., standard combination may include a macrolide (clarithromycin, azithromycin), a rifamycin (rifampin, rifabutin), and ethambutol] AND
- 3. Patient will continue treatment with a combination antibiotic therapy for MAC lung disease with the requested agent [e.g., combination may include a macrolide (clarithromycin, azithromycin), a rifamycin (rifampin, rifabutin), and ethambutol]

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Mycobacterium avium complex (MAC) lung disease AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will continue treatment with a combination antibiotic therapy for MAC lung disease with the requested agent [e.g., combination may include a macrolide (clarithromycin, azithromycin), a rifamycin (rifampin, rifabutin), and ethambutol]

# Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., infectious disease, immunologist, pulmonologist, thoracic specialist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Armodafinil PA

# Drug Name(s)

Armodafinil

Nuvigil

# Indications:

All Medically-Accepted Indications.

# Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another target agent (i.e., modafinil)

# Age Restriction:

Patient is 17 years of age or over

# **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

Asceniv PA

# Drug Name(s)

Asceniv

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for approval require ONE of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Primary immunodeficiency [e.g., congenital agammaglobulinemia, common variable immunodeficiency (CVID), severe combined immunodeficiency, Wiskott-Aldrich Syndrome, X-linked agammaglobulinemia (XLA), humoral immunodeficiency, IgG subclass deficiency with or without IgA deficiency] OR
  - B. B-cell chronic lymphocytic leukemia OR multiple myeloma AND ONE of the following:
    - i. Patient has a history of infections OR
    - ii. Patient has evidence of specific antibody deficiency OR
    - iii. Patient has hypogammaglobulinemia OR
  - C. Idiopathic thrombocytopenia purpura AND ONE of the following:
    - i. Patient has failed ONE conventional therapy [e.g., corticosteroids (e.g., methylprednisolone), or immunosuppressants (e.g., azathioprine)] OR
    - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to ONE conventional therapy OR
  - D. Dermatomyositis AND ONE of the following:
    - i. Patient has failed ONE conventional therapy [e.g., corticosteroids (e.g., methylprednisolone) or immunosuppressants (e.g., azathioprine)] OR
    - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to ONE conventional therapy OR
  - E. Polymyositis AND ONE of the following:
    - i. Patient has failed ONE conventional therapy [e.g., corticosteroids (e.g., methylprednisolone) or immunosuppressants (e.g., azathioprine)] OR
    - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to ONE conventional therapy OR
  - F. Severe rheumatoid arthritis AND ONE of the following:
    - i. Patient has failed ONE conventional therapy [e.g., tumor necrosis factor antagonists (e.g., Enbrel), DMARDS (e.g., methotrexate)] OR
    - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to ONE conventional therapy OR

Criteria continues: see Other Criteria

Age Restriction:

Prescriber Restrictions: Coverage Duration:

Approval will be for 6 months for indications in Other Criteria, 12 months for all others **Other Criteria**:

- G. Myasthenia gravis (MG) AND ONE of the following:
  - i. Patient is in acute myasthenic crisis OR
  - ii. Patient has severe refractory MG (e.g., major functional disability/weakness) AND ONE of the following:
    - a) Patient has failed ONE immunomodulator therapy (i.e., corticosteroid, pyridostigmine, or azathioprine) OR
    - b) Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to ONE immunomodulator therapy OR
- H. Multiple sclerosis (MS) AND BOTH of the following:
  - i. Patient has a diagnosis of relapsing remitting MS (RRMS) AND
  - ii. Patient has had an insufficient response, documented failure, or FDA labeled contraindication to TWO MS agents (e.g., Avonex, Betaseron, Copaxone, dimethyl fumarate, fingolimod, glatiramer, Glatopa, Kesimpta, Plegridy, teriflunomide, Vumerity) OR
- I. Acquired von Willebrand hemophilia AND ONE of the following:
  - i. Patient has failed ONE conventional therapy (e.g., desmopressin, von Willebrand factor replacement therapy, corticosteroids, cyclophosphamide, FEIBA, or recombinant factor VIIa) OR
  - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to ONE conventional therapy OR
- J. Refractory pemphigus vulgaris AND ONE of the following:
  - i. Patient has failed ONE conventional immunosuppressive therapy (e.g., azathioprine, cyclophosphamide, mycophenolate, corticosteroids) OR
  - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to ONE conventional immunosuppressive therapy OR
- 2. ONE of the following:
  - A. Patient has another FDA labeled indication for the requested agent OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

Indications with 6 months approval duration: Acquired von Willebrand hemophilia, Guillain-Barre Syndrome, Lambert-Eaton myasthenia syndrome, Kawasaki disease, CMV induced pneumonitis in solid organ transplant, Toxic shock syndrome due to invasive group A streptococcus, Toxic epidermal necrolysis and Stevens-Johnson syndrome

Drug is also subject to Part B versus Part D review.

Atopic Dermatitis PA – Eucrisa

# Drug Name(s)

Eucrisa

#### **Indications:**

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of atopic dermatitis AND
- 2. ONE of the following:
  - A. Patient has tried and had an inadequate response to a topical corticosteroid or topical corticosteroid combination preparation (e.g., hydrocortisone, triamcinolone) OR
  - B. Patient has an intolerance or hypersensitivity to a topical corticosteroid or topical corticosteroid combination preparation OR
  - C. Patient has an FDA labeled contraindication to a topical corticosteroid or topical corticosteroid combination preparation

**Age Restriction:** 

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Atopic Dermatitis PA – Pimecrolimus

# Drug Name(s)

Elidel

Pimecrolimus

#### **Indications:**

All Medically-Accepted Indications.

#### **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require ONE of the following:

- 1. Patient has a diagnosis of atopic dermatitis or vulvar lichen sclerosus AND ONE of the following:
  - A. Patient has tried and had an inadequate response to a topical corticosteroid or topical corticosteroid combination preparation (e.g., hydrocortisone, triamcinolone) OR
  - B. Patient has an intolerance or hypersensitivity to a topical corticosteroid or topical corticosteroid combination preparation OR
  - C. Patient has an FDA labeled contraindication to a topical corticosteroid or topical corticosteroid combination preparation OR
- 2. Patient has a diagnosis of facial seborrheic dermatitis associated with HIV infection AND BOTH of the following:
  - A. Patient is currently on an antiretroviral treatment regimen AND
  - B. ONE of the following:
    - i. Patient has tried and had an inadequate response to a topical corticosteroid or topical antifungal treatment (e.g., hydrocortisone, triamcinolone, ketoconazole) OR
    - ii. Patient has an intolerance or hypersensitivity to a topical corticosteroid or topical antifungal treatment OR
    - iii. Patient has an FDA labeled contraindication to a topical corticosteroid or topical antifungal treatment OR
- 3. Patient has an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Atopic Dermatitis PA – Tacrolimus

# Drug Name(s)

Tacrolimus

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require ONE of the following:

- 1. Patient has a diagnosis of atopic dermatitis AND ONE of the following:
  - A. Patient has tried and had an inadequate response to a topical corticosteroid or topical corticosteroid combination preparation (e.g., hydrocortisone, triamcinolone) OR
  - B. Patient has an intolerance or hypersensitivity to a topical corticosteroid or topical corticosteroid combination preparation OR
  - C. Patient has an FDA labeled contraindication to a topical corticosteroid or topical corticosteroid combination preparation OR
- 2. Patient has an indication that is supported in CMS approved compendia for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Atovaquone PA

# Drug Name(s)

Atovaquone

Mepron

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - 1. Patient has a diagnosis of mild-to-moderate Pneumocystis jirovecii pneumonia OR
      - 2. Patient is using the requested agent for prevention of Pneumocystis jirovecii pneumonia AND
    - ii. ONE of the following:
      - 1. Patient has an intolerance or hypersensitivity to trimethoprim/sulfamethoxazole (TMP/SMX) OR
      - 2. Patient has an FDA labeled contraindication to trimethoprim/sulfamethoxazole (TMP/SMX) OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Austedo PA

# Drug Name(s)

Austedo

Austedo Xr

Austedo Xr Titration Kit

### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chorea associated with Huntington's disease AND BOTH of the following:
    - i. ONE of the following:
      - 1. Patient does NOT have a current diagnosis of depression OR
      - 2. Patient has a current diagnosis of depression and is being treated for depression AND
    - ii. ONE of the following:
      - 1. Patient does NOT have a diagnosis of passive suicidal ideation and/or behavior OR
      - 2. Patient has a diagnosis of passive suicidal ideation and/or behavior and must NOT be actively suicidal OR
  - B. Patient has a diagnosis of tardive dyskinesia AND ONE of the following:
    - i. Prescriber has reduced the dose of or discontinued any medications known to cause tardive dyskinesia (i.e., dopamine receptor blocking agents) OR
    - ii. Prescriber has provided clinical rationale indicating that a reduced dose or discontinuation of any medications known to cause tardive dyskinesia is not appropriate AND
- 2. Patient will NOT be using the requested agent in combination with a monoamine oxidase inhibitor (MAOI) AND
- 3. Patient will NOT be using the requested agent in combination with reserpine

#### Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Azstarys PA

Drug Name(s)

Azstarys

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Belsomra PA

Drug Name(s)

Belsomra

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Benign Prostatic Hyperplasia PA – Tadalafil

# Drug Name(s)

Cialis

Tadalafil 2.5Mg, 5Mg

### **Indications:**

All FDA-Approved Indications.

# Off-Label Uses:

### **Exclusion Criteria:**

Requested agent will be used to treat erectile dysfunction AND FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of benign prostatic hyperplasia (BPH) AND
- 2. Patient has tried and had an insufficient response, intolerance or hypersensitivity, or FDA labeled contraindication to TWO alpha blocker agents (e.g., terazosin, doxazosin, tamsulosin)

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Benlysta SC PA

Drug Name(s)

Benlysta SC

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Benzodiazepines PA - Chlordiazepoxide

# Drug Name(s)

Chlordiazepoxide Hcl

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. BOTH of the following:
    - i. Patient has ONE of the following diagnoses:
      - a. Anxiety disorder AND ONE of the following:
        - 1) Patient has tried and had an inadequate response to a formulary selective serotonin reuptake inhibitor (SSRI) or serotonin norepinephrine reuptake inhibitor (SNRI) OR
        - 2) Patient has an intolerance or hypersensitivity to a formulary SSRI or SNRI OR
        - 3) Patient has an FDA labeled contraindication to a formulary SSRI or SNRI OR
      - b. Alcohol withdrawal OR
      - c. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Benzodiazepines PA – Chlordiazepoxide /amitriptyline

### Drug Name(s)

Chlordiazepoxide/Amitriptyline

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. BOTH of the following:
    - i. Patient has ONE of the following diagnoses:
      - a. Moderate to severe depression with moderate to severe anxiety AND ONE of the following:
        - 1) Patient has tried and had an inadequate response to a formulary selective serotonin reuptake inhibitor (SSRI) or serotonin norepinephrine reuptake inhibitor (SNRI) OR
        - 2) Patient has an intolerance or hypersensitivity to a formulary SSRI or SNRI OR
        - 3) Patient has an FDA labeled contraindication to a formulary SSRI or SNRI OR
      - b. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Benzodiazepines PA - Clobazam

# Drug Name(s)

Clobazam

Onfi

#### **Indications:**

All Medically-Accepted Indications.

#### **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. BOTH of the following:
    - i. Patient has ONE of the following diagnoses:
      - a. Seizure disorder OR
      - b. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Benzodiazepines PA – Clorazepate

# Drug Name(s)

Clorazepate Dipotassium

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. BOTH of the following:
    - i. Patient has ONE of the following diagnoses:
      - a. Seizure disorder OR
      - b. Anxiety disorder AND ONE of the following:
        - 1) Patient has tried and has an inadequate response to a formulary selective serotonin reuptake inhibitor (SSRI) or serotonin norepinephrine reuptake inhibitor (SNRI) OR
        - 2) Patient has an intolerance or hypersensitivity to a formulary SSRI or SNRI OR
        - 3) Patient has an FDA labeled contraindication to a formulary SSRI or SNRI OR
      - c. Alcohol withdrawal OR
      - d. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent

### **Age Restriction:**

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Benzodiazepines PA - Diazepam

# Drug Name(s)

Diazepam

Diazepam Intensol

Valium

### **Indications:**

All Medically-Accepted Indications.

### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. BOTH of the following:
    - i. Patient has ONE of the following diagnoses:
      - a. Seizure disorder OR
      - b. Anxiety disorder AND ONE of the following:
        - 1) Patient has tried and had an inadequate response to a formulary selective serotonin reuptake inhibitor (SSRI) or serotonin norepinephrine reuptake inhibitor (SNRI) OR
        - 2) Patient has an intolerance or hypersensitivity to a formulary SSRI or SNRI OR
        - 3) Patient has an FDA labeled contraindication to a formulary SSRI or SNRI OR
      - c. Skeletal muscle spasms OR
      - d. Alcohol withdrawal OR
      - e. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Benzodiazepines PA – Estazolam

Drug Name(s)

Estazolam

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require the following:

1. Patient has a diagnosis of insomnia

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Benzodiazepines PA – Flurazepam

# Drug Name(s)

Flurazepam Hcl

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require the following:

1. Patient has a diagnosis of insomnia

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Benzodiazepines PA – Lorazepam

# Drug Name(s)

Ativan

Lorazepam

Lorazepam Intensol

Loreev Xr

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. BOTH of the following:
    - i. Patient has ONE of the following diagnoses:
      - a. Anxiety disorder AND ONE of the following:
        - 1) Patient has tried and had an inadequate response to a formulary selective serotonin reuptake inhibitor (SSRI) or serotonin norepinephrine reuptake inhibitor (SNRI) OR
        - 2) Patient has an intolerance or hypersensitivity to a formulary SSRI or SNRI OR
        - 3) Patient has an FDA labeled contraindication to a formulary SSRI or SNRI OR
      - b. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Benzodiazepines PA – Oxazepam

## Drug Name(s)

Oxazepam

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. BOTH of the following:
    - i. Patient has ONE of the following diagnoses:
      - a. Anxiety disorder AND ONE of the following:
        - 1) Patient has tried and had an inadequate response to a formulary selective serotonin reuptake inhibitor (SSRI) or serotonin norepinephrine reuptake inhibitor (SNRI) OR
        - 2) Patient has an intolerance or hypersensitivity to a formulary SSRI or SNRI OR
        - 3) Patient has an FDA labeled contraindication to a formulary SSRI or SNRI OR
      - b. Alcohol withdrawal OR
      - c. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

Benzodiazepines PA – Sympazan

# Drug Name(s)

Sympazan

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. BOTH of the following:
    - i. Patient has ONE of the following diagnoses:
      - a. Seizure disorder OR
      - b. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

Bethkis PA

Drug Name(s)

Bethkis

Tobramycin (Bethkis)

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Bexarotene Gel PA

## Drug Name(s)

Bexarotene Gel

Targretin Gel

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. ALL of the following:
    - i. ONE of the following:
      - 1. BOTH of the following:
        - a. Patient has a diagnosis of stage IA or IB cutaneous T-cell lymphoma (CTCL) with cutaneous lesions AND
        - b. ONE of the following:
          - i. Patient has refractory or persistent disease despite a previous treatment trial with a skin-directed therapy (e.g., topical corticosteroid, topical imiquimod) OR
          - ii. Patient has an intolerance or hypersensitivity to a previous treatment trial with a skin-directed therapy (e.g., topical corticosteroid, topical imiquimod) OR
          - iii. Patient has an FDA labeled contraindication to a previous treatment trial with a skin-directed therapy (e.g., topical corticosteroid, topical imiquimod) OR
      - 2. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Prescriber is a specialist in the area of the patient's diagnosis (e.g., dermatologist, oncologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis AND
    - iii. Patient does NOT have any FDA labeled contraindications to the requested agent

## Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 3. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. ALL of the following:
    - i. Patient has had clinical benefit with the requested agent AND
    - ii. Prescriber is a specialist in the area of the patient's diagnosis (e.g., dermatologist, oncologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis AND
    - iii. Patient does NOT have any FDA labeled contraindications to the requested agent

Biologic Immunomodulators PA – Abrilada

Drug Name(s)

Abrilada

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Pending CMS Review

Biologic Immunomodulators PA – Actemra

# Drug Name(s)

Actemra

Actemra Actpen

**Indications:** 

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Pending CMS Review

Biologic Immunomodulators PA – Amjevita

## Drug Name(s)

Amjevita

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

## Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of juvenile idiopathic arthritis

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of ulcerative colitis

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of pediatric Crohn's disease

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Use of TWO preferred agents (Cosentyx, Hadlima, or Humira) is required for diagnosis of hidradenitis suppurativa

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of uveitis

Biologic Immunomodulators PA – Bimzelx

## Drug Name(s)

Bimzelx

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Other Criteria:

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Biologic Immunomodulators PA - Cimzia

## Drug Name(s)

Cimzia

Cimzia Starter Kit

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of Crohn's disease

Only the preferred agent Cosentyx is required for diagnosis of non-radiographic axial spondyloarthritis

Biologic Immunomodulators PA - Cosentyx

## Drug Name(s)

Cosentyx

Cosentyx Sensoready Pen

Cosentyx Unoready

#### **Indications:**

All FDA-Approved Indications.

## Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's medication history indicates use of another biologic immunomodulator agent for the same FDA labeled indication OR
  - D. Patient's diagnosis does NOT require a conventional prerequisite agent OR
  - E. Patient's medication history indicates use of ONE formulary conventional prerequisite agent for the requested indication OR
  - F. Patient has an intolerance or hypersensitivity to at least ONE formulary conventional prerequisite agent for the requested indication OR
  - G. Patient has an FDA labeled contraindication to at least ONE formulary conventional prerequisite agent for the requested indication AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

# Other Criteria:

Use of ONE conventional prerequisite agent is required for diagnoses of psoriatic arthritis or plaque psoriasis

NO prerequisites are required for diagnoses of ankylosing spondylitis, enthesitis related arthritis, hidradenitis suppurativa, or non-radiographic axial spondyloarthritis

Formulary conventional agents for psoriatic arthritis include cyclosporine, leflunomide, methotrexate, or sulfasalazine

Formulary conventional agents (topical or systemic) for plaque psoriasis include acitretin, calcipotriene, methotrexate, tazarotene, or topical corticosteroids

Biologic Immunomodulators PA - Cyltezo

## Drug Name(s)

Cyltezo

Cyltezo Starter Package For Crohns Disease/Uc/Hs

Cyltezo Starter Package For Psoriasis

Cyltezo Starter Package For Psoriasis/Uveitis Indications:

All FDA-Approved Indications.

#### **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

## **Prescriber Restrictions:**

## **Coverage Duration:**

Approval will be 12 weeks for initial use for ulcerative colitis, 12 months for all others

#### Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of juvenile idiopathic arthritis

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of ulcerative colitis

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of pediatric Crohn's disease

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Use of TWO preferred agents (Cosentyx, Hadlima, or Humira) is required for diagnosis of hidradenitis suppurativa

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of uveitis

Biologic Immunomodulators PA - Enbrel

# Drug Name(s)

**Enbrel** 

Enbrel Mini

**Enbrel Sureclick** 

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's medication history indicates use of another biologic immunomodulator agent for the same FDA labeled indication OR
  - D. Patient's diagnosis does NOT require a conventional prerequisite agent OR
  - E. Patient's medication history indicates use of ONE formulary conventional prerequisite agent for the requested indication OR
  - F. Patient has an intolerance or hypersensitivity to at least ONE formulary conventional prerequisite agent for the requested indication OR
  - G. Patient has an FDA labeled contraindication to at least ONE formulary conventional prerequisite agent for the requested indication AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

# Other Criteria:

Use of ONE conventional prerequisite agent is required for diagnoses of psoriatic arthritis, plaque psoriasis, rheumatoid arthritis, juvenile psoriatic arthritis, or juvenile idiopathic arthritis

NO prerequisites are required for a diagnoses of ankylosing spondylitis or severe juvenile psoriatic arthritis

Formulary conventional agents for rheumatoid arthritis, juvenile idiopathic arthritis, juvenile psoriatic arthritis, or psoriatic arthritis include leflunomide, methotrexate, or sulfasalazine

Formulary conventional agents (topical or systemic) for plaque psoriasis include acitretin, calcipotriene, methotrexate, tazarotene, or topical corticosteroids

Biologic Immunomodulators PA – Entyvio SC

# Drug Name(s)

Entyvio Pen

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Pending CMS Review

Biologic Immunomodulators PA – Hadlima

## Drug Name(s)

Hadlima

Hadlima Pushtouch

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's medication history indicates use of another biologic immunomodulator agent for the same FDA labeled indication OR
  - D. Patient's diagnosis does NOT require a conventional prerequisite agent OR
  - E. Patient's medication history indicates use of ONE formulary conventional prerequisite agent for the requested indication OR
  - F. Patient has an intolerance or hypersensitivity to at least ONE formulary conventional prerequisite agent for the requested indication OR
  - G. Patient has an FDA labeled contraindication to at least ONE formulary conventional prerequisite agent for the requested indication AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

## **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be 12 weeks for initial use for ulcerative colitis, 12 months for all others

#### Other Criteria:

Use of ONE conventional prerequisite agent is required for diagnoses of psoriatic arthritis, plaque psoriasis, rheumatoid arthritis, juvenile idiopathic arthritis, Crohn's disease, or moderate ulcerative colitis

NO prerequisites are required for diagnoses of ankylosing spondylitis, hidradenitis suppurativa, severe ulcerative colitis, or uveitis

Formulary conventional agents for rheumatoid arthritis, juvenile idiopathic arthritis, or psoriatic arthritis include leflunomide, methotrexate, or sulfasalazine

Formulary conventional agents (topical or systemic) for plaque psoriasis include acitretin, calcipotriene, methotrexate, tazarotene, or topical corticosteroids

Formulary conventional agents for Crohn's disease include methotrexate, sulfasalazine, corticosteroids, azathioprine, or mercaptopurine

Formulary conventional agents for moderate ulcerative colitis include 5-aminosalicylates, corticosteroids, azathioprine, or mercaptopurine

Biologic Immunomodulators PA - Hulio

## Drug Name(s)

Adalimumab-Fkjp

Hulio

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

## **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of juvenile idiopathic arthritis

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of ulcerative colitis

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of pediatric Crohn's disease

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Use of TWO preferred agents (Cosentyx, Hadlima, or Humira) is required for diagnosis of hidradenitis suppurativa

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of uveitis

Biologic Immunomodulators PA - Humira

# Drug Name(s)

Humira

Humira Pen

Humira Pen-Cd/Uc/Hs Starter

Humira Pen-Pediatric Uc Starter Pack

Humira Pen-Ps/Uv Starter

#### Indications:

All FDA-Approved Indications.

#### **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's medication history indicates use of another biologic immunomodulator agent for the same FDA labeled indication OR
  - D. Patient's diagnosis does NOT require a conventional prerequisite agent OR
  - E. Patient's medication history indicates use of ONE formulary conventional prerequisite agent for the requested indication OR
  - F. Patient has an intolerance or hypersensitivity to at least ONE formulary conventional prerequisite agent for the requested indication OR
  - G. Patient has an FDA labeled contraindication to at least ONE formulary conventional prerequisite agent for the requested indication AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

## **Prescriber Restrictions:**

## **Coverage Duration:**

Approval will be 12 weeks for initial use for ulcerative colitis, 12 months for all others

#### Other Criteria:

Use of ONE conventional prerequisite agent is required for diagnoses of psoriatic arthritis, plaque psoriasis, rheumatoid arthritis, juvenile idiopathic arthritis, Crohn's disease, or moderate ulcerative colitis

NO prerequisites are required for diagnoses of ankylosing spondylitis, hidradenitis suppurativa, severe ulcerative colitis, or uveitis

Formulary conventional agents for rheumatoid arthritis, juvenile idiopathic arthritis, or psoriatic arthritis include leflunomide, methotrexate, or sulfasalazine

Formulary conventional agents (topical or systemic) for plaque psoriasis include acitretin, calcipotriene, methotrexate, tazarotene, or topical corticosteroids

Formulary conventional agents for Crohn's disease include methotrexate, sulfasalazine, corticosteroids, azathioprine, or mercaptopurine

Formulary conventional agents for moderate ulcerative colitis include 5-aminosalicylates, corticosteroids, azathioprine, or mercaptopurine

Biologic Immunomodulators PA – Hyrimoz

## Drug Name(s)

Adalimumab-Adaz

Hyrimoz

Hyrimoz Crohns Disease And Ulcerative Colitis Starter Pack

Hyrimoz Pediatric Crohns Disease Starter Pack

Hyrimoz Plaque Psoriasis/Uveitis Starter Pack

Hyrimoz Sensoready Pen

#### **Indications:**

All FDA-Approved Indications.

## Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND

- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of juvenile idiopathic arthritis

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of ulcerative colitis

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of pediatric Crohn's disease

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Use of TWO preferred agents (Cosentyx, Hadlima, or Humira) is required for diagnosis of hidradenitis suppurativa

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of uveitis

Biologic Immunomodulators PA - Idacio

## Drug Name(s)

Adalimumab-Aacf (2 Pen)

Adalimumab-Aacf (2 Syringe)

Adalimumab-Aacf Starter Pack/Cd/Uc/Hs (6 Pen)

Adalimumab-Aacf Starter Pack/Psoriasis/Uveitis (4 Pen)

Idacio (2 Pen)

Idacio (2 Syringe)

Idacio Starter Package For Crohns Disease

Idacio Starter Package For Plaque Psoriasis

## **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND

- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator

#### Age Restriction:

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of juvenile idiopathic arthritis

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of ulcerative colitis

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of pediatric Crohn's disease

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Use of TWO preferred agents (Cosentyx, Hadlima, or Humira) is required for diagnosis of hidradenitis suppurativa

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of uveitis

Biologic Immunomodulators PA – Ilumya

## Drug Name(s)

Ilumya

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Other Criteria:

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Biologic Immunomodulators PA - Inflectra

## Drug Name(s)

Inflectra

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

## Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of adult ulcerative colitis

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of pediatric Crohn's disease

Only the preferred agent Humira is required for diagnosis of pediatric ulcerative colitis

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Biologic Immunomodulators PA – Kevzara

## Drug Name(s)

Kevzara

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

# Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of rheumatoid arthritis

NO preferred agent is required for diagnosis of polymyalgia rheumatica

Biologic Immunomodulators PA - Kineret

# Drug Name(s)

Kineret

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

### **Prescriber Restrictions:**

### **Coverage Duration:**

Approval will be for 12 months

# Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of rheumatoid arthritis

NO preferred agent is required for diagnoses of Neonatal-Onset Multisystem Inflammatory Disease (NOMID) or Deficiency of Interleukin-1 Receptor Antagonist (DIRA)

Biologic Immunomodulators PA - Litfulo

# Drug Name(s)

Litfulo

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Biologic Immunomodulators PA – Olumiant

# Drug Name(s)

Olumiant

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

### **Prescriber Restrictions:**

### **Coverage Duration:**

Approval will be for 12 months

# Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of rheumatoid arthritis

NO preferred agent is required for diagnosis of alopecia areata

Biologic Immunomodulators PA - Omvoh

# Drug Name(s)

Omvoh

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

e of TWO preferred agents (Hadlima, Humira, Rinvoq tablets, or Stelara) is required for diagnosis of erative colitis	of

Biologic Immunomodulators PA - Orencia

## Drug Name(s)

Orencia

Orencia Clickject

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

# **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, Humira, Rinvoq tablets, or Rinvoq solution) is required for diagnosis of juvenile idiopathic arthritis

Use of TWO preferred agents (Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of rheumatoid arthritis

For patients 18 years of age or over, use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Rinvoq tablets, Rinvoq solution, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

For patients between 6 and less than 18 years of age, use of TWO preferred agents (Cosentyx, Rinvoq tablets, or Rinvoq solution) is required for diagnosis of psoriatic arthritis

For patients between 2 and less than 6 years of age, use of ONE preferred agent (Rinvoq tablets or Rinvoq solution) is required for diagnosis of psoriatic arthritis

NO preferred agent is required for diagnosis of prophylaxis of acute graft vs host disease

Biologic Immunomodulators PA – Remicade

## Drug Name(s)

Remicade

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

### **Prescriber Restrictions:**

### **Coverage Duration:**

Approval will be for 12 months

# Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of adult ulcerative colitis

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of pediatric Crohn's disease

Only the preferred agent Humira is required for diagnosis of pediatric ulcerative colitis

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Biologic Immunomodulators PA – Renflexis

## Drug Name(s)

Renflexis

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

### **Prescriber Restrictions:**

### **Coverage Duration:**

Approval will be for 12 months

# Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of adult ulcerative colitis

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of pediatric Crohn's disease

Only the preferred agent Humira is required for diagnosis of pediatric ulcerative colitis

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Biologic Immunomodulators PA - Riabni

## Drug Name(s)

Riabni

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - ii. Prescriber states the patient is currently being treated with the requested agent OR B. ALL of the following:
    - i. ONE of the following:
      - a. Patient has a diagnosis of rheumatoid arthritis AND ONE of the following:
        - 1. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
        - 2. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
        - 3. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
      - b. Patient has another FDA labeled indication or an indication that is supported in CMS approved compendia AND
    - ii. Patient has been screened for hepatitis B infection measuring hepatitis B surface antigen (HBsAg) and hepatitis B core antibody (anti-HBc) and has begun therapy, if appropriate, prior to receiving the requested agent AND
    - iii. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
    - iv. Patient does NOT have any FDA labeled limitation(s) of use that is not otherwise supported in NCCN guidelines AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

### Age Restriction:

### **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

# Other Criteria:

Criteria for renewal approval require ALL of the following:

1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND

- 2. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 3. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. ALL of the following:
    - i. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
    - ii. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
    - iii. Patient does NOT have any FDA labeled limitation(s) of use that is not otherwise supported in NCCN guidelines AND
- 4. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Use of TWO preferred agents (Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of rheumatoid arthritis

ALL other diagnoses do NOT require any preferred agents

Biologic Immunomodulators PA – Rinvoq Solution

## Drug Name(s)

Rinvoq Lq

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's medication history indicates use of preferred TNF agent(s) OR
    - ii. Patient has an intolerance or hypersensitivity to preferred TNF agent(s) OR
    - iii. Patient has an FDA labeled contraindication to preferred TNF agent(s) OR
    - iv. The request is for an FDA labeled indication that is not covered by preferred TNF agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

### Age Restriction:

### **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Use of ONE preferred TNF (Enbrel, Hadlima, or Humira) is required for diagnoses of adult psoriatic arthritis or juvenile idiopathic arthritis

NO preferred TNF agent is required for diagnosis of pediatric psoriatic arthritis

Biologic Immunomodulators PA – Rinvoq Tablet

## Drug Name(s)

Rinvoq Tablet

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. BOTH of the following:
      - a. Patient has an FDA labeled indication other than moderate to severe atopic dermatitis for the requested agent AND
      - b. ONE of the following:
        - 1. Patient's medication history indicates use of preferred TNF agent(s) OR
        - 2. Patient has an intolerance or hypersensitivity to preferred TNF agent(s) OR
        - 3. Patient has an FDA labeled contraindication to preferred TNF agent(s) OR
        - 4. The request is for an FDA labeled indication that is not covered by preferred TNF agent(s) OR
    - ii. Patient has a diagnosis of moderate to severe atopic dermatitis AND ONE of the following:
      - a. Patient's medication history indicates use of TWO conventional prerequisite agents (i.e., ONE formulary topical corticosteroid AND ONE formulary topical calcineurin inhibitor) for the requested indication OR
      - b. Patient has an intolerance or hypersensitivity to TWO conventional prerequisite agents (i.e., ONE formulary topical corticosteroid AND ONE formulary topical calcineurin inhibitor) for the requested indication OR
      - c. Patient has an FDA labeled contraindication to TWO conventional prerequisite agents (i.e., ONE formulary topical corticosteroid AND ONE formulary topical calcineurin inhibitor) for the requested indication AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Age Restriction:

**Prescriber Restrictions:** 

## **Coverage Duration:**

Approval will be for 12 months

### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

Use of ONE preferred TNF (Enbrel, Hadlima, or Humira) is required for diagnoses of ankylosing spondylitis, rheumatoid arthritis, adult psoriatic arthritis, or juvenile idiopathic arthritis

Use of ONE preferred TNF (Hadlima or Humira) is required for diagnoses of ulcerative colitis or Crohn's disease

Use of TWO conventional prerequisite agents are required for diagnosis of moderate to severe atopic dermatitis

NO preferred TNF agents are required for diagnoses of pediatric psoriatic arthritis or non-radiographic axial spondyloarthritis

Biologic Immunomodulators PA - Ruxience

## Drug Name(s)

Ruxience

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - ii. Prescriber states the patient is currently being treated with the requested agent OR B. ALL of the following:
    - i. ONE of the following:
      - a. Patient has a diagnosis of rheumatoid arthritis AND ONE of the following:
        - 1. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
        - 2. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
        - 3. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
      - b. Patient has another FDA labeled indication or an indication that is supported in CMS approved compendia AND
    - ii. Patient has been screened for hepatitis B infection measuring hepatitis B surface antigen (HBsAg) and hepatitis B core antibody (anti-HBc) and has begun therapy, if appropriate, prior to receiving the requested agent AND
    - iii. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
    - iv. Patient does NOT have any FDA labeled limitation(s) of use that is not otherwise supported in NCCN guidelines AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

# Other Criteria:

Criteria for renewal approval require ALL of the following:

1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND

- 2. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 3. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. ALL of the following:
    - i. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
    - ii. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
    - iii. Patient does NOT have any FDA labeled limitation(s) of use that is not otherwise supported in NCCN guidelines AND
- 4. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Use of TWO preferred agents (Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of rheumatoid arthritis

ALL other diagnoses do NOT require any preferred agents

Biologic Immunomodulators PA - Siliq

## Drug Name(s)

Siliq

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Biologic Immunomodulators PA - Simlandi

# Drug Name(s)

Simlandi 1Pn Kit

Simlandi 2Pn Inj

# **Indications:**

All FDA-Approved Indications

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Pending CMS Review

Biologic Immunomodulators PA - Simponi

## Drug Name(s)

Simponi

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of ulcerative colitis

Biologic Immunomodulators PA - Skyrizi

# Drug Name(s)

Skyrizi

Skyrizi Pen

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's medication history indicates use of another biologic immunomodulator agent for the same FDA labeled indication OR
  - D. Patient's medication history indicates use of ONE formulary conventional prerequisite agent for the requested indication OR
  - E. Patient has an intolerance or hypersensitivity to at least ONE formulary conventional prerequisite agent for the requested indication OR
  - F. Patient has an FDA labeled contraindication to at least ONE formulary conventional prerequisite agent for the requested indication AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

**Prescriber Restrictions:** 

## **Coverage Duration:**

Approval will be for 12 months

Use of ONE conventional prerequisite agent is required for diagnoses of Crohn's disease, plaque psoriasis, or psoriatic arthritis

Formulary conventional agents for Crohn's disease include methotrexate, sulfasalazine, corticosteroids, azathioprine, or mercaptopurine

Formulary conventional agents (topical or systemic) for plaque psoriasis include acitretin, calcipotriene, methotrexate, tazarotene, or topical corticosteroids

Formulary conventional agents for psoriatic arthritis include leflunomide, methotrexate, or sulfasalazine

Biologic Immunomodulators PA – Sotyktu

# Drug Name(s)

Sotyktu

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Biologic Immunomodulators PA - Stelara

# Drug Name(s)

Stelara

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's medication history indicates use of another biologic immunomodulator agent for the same FDA labeled indication OR
  - D. Patient's diagnosis does NOT require a conventional prerequisite agent OR
  - E. Patient's medication history indicates use of ONE formulary conventional prerequisite agent for the requested indication OR
  - F. Patient has an intolerance or hypersensitivity to at least ONE formulary conventional prerequisite agent for the requested indication OR
  - G. Patient has an FDA labeled contraindication to at least ONE formulary conventional prerequisite agent for the requested indication AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Use of ONE conventional prerequisite agent is required for diagnoses of psoriatic arthritis, plaque psoriasis, moderate ulcerative colitis, or Crohn's disease

NO prerequisites are required for diagnosis of severe ulcerative colitis

Formulary conventional agents for psoriatic arthritis include cyclosporine, leflunomide, methotrexate, or sulfasalazine

Formulary conventional agents (topical or systemic) for plaque psoriasis include acitretin, calcipotriene, methotrexate, tazarotene, or topical corticosteroids

Formulary conventional agents for Crohn's disease include methotrexate, sulfasalazine, corticosteroids, azathioprine, mercaptopurine

Formulary conventional agents for moderate ulcerative colitis include 5-aminosalicylates, corticosteroids, azathioprine, mercaptopurine

Biologic Immunomodulators PA - Taltz

# Drug Name(s)

Talt<sub>7</sub>

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Rinvoq tablets, Rinvoq solution, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

For patients 18 years of age or over, use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

For patients between 6 and less than 18 years of age, use of TWO preferred agents (Cosentyx, Enbrel, or Stelara) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Cosentyx and Rinvoq tablets) is required for diagnosis of non-radiographic axial spondyloarthritis

Biologic Immunomodulators PA - Tremfya

## Drug Name(s)

Tremfya

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's medication history indicates use of another biologic immunomodulator agent for the same FDA labeled indication OR
  - D. Patient's medication history indicates use of ONE formulary conventional prerequisite agent for the requested indication OR
  - E. Patient has an intolerance or hypersensitivity to at least ONE formulary conventional prerequisite agent for the requested indication OR
  - F. Patient has an FDA labeled contraindication to at least ONE formulary conventional prerequisite agent for the requested indication AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Use of ONE conventional prerequisite agent is required for diagnoses of psoriatic arthritis or plaque psoriasis

Formulary conventional agents for psoriatic arthritis include cyclosporine, leflunomide, methotrexate, or sulfasalazine

Formulary conventional agents (topical or systemic) for plaque psoriasis include acitretin, calcipotriene, methotrexate, tazarotene, or topical corticosteroids

Biologic Immunomodulators PA - Tyenne

# Drug Name(s)

Tyenne Inj

Indications:

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Pending CMS Review

Biologic Immunomodulators PA - Velsipity

### Drug Name(s)

Velsipity

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Use of TWO preferred agents (Hadlima, Humira, Rinvoq tablets, or Stelara) is required for diagnosis of ulcerative colitis

Biologic Immunomodulators PA - Xeljanz Solution

# Drug Name(s)

Xeljanz Solution

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Use of TWO preferred agents (Enbrel, Hadlima, Humira, Rinvoq tablets, or Rinvoq solution) is required for diagnosis of juvenile idiopathic arthritis

Biologic Immunomodulators PA - Xeljanz Tablet

### Drug Name(s)

Xeljanz Tablet

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Rinvoq tablets, Rinvoq solution, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Enbrel, Hadlima, Humira, Rinvoq tablets, or Rinvoq solution) is required for diagnosis of juvenile idiopathic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Rinvoq tablets, or Stelara) is required for diagnosis of ulcerative colitis

Biologic Immunomodulators PA - Xeljanz XR

### Drug Name(s)

Xeljanz Xr

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - C. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
    - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
    - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
    - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Rinvoq tablets, Rinvoq solution, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, or Rinvoq tablets) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Rinvoq tablets, or Stelara) is required for diagnosis of ulcerative colitis

Biologic Immunomodulators PA - Yuflyma

# Drug Name(s)

Adalimumab-Aaty 1-Pen Kit

Adalimumab-Aaty 2-Pen Kit

Adalimumab-Aaty 2-Syringe Kit

Yuflyma 1-Pen Kit

Yuflyma 2-Pen Kit

Yuflyma 2-Syringe Kit

Yuflyma Cd/Uc/Hs Starter

#### **Indications:**

All FDA-Approved Indications.

#### **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND

- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

#### **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of juvenile idiopathic arthritis

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of ulcerative colitis

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of pediatric Crohn's disease

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Use of TWO preferred agents (Cosentyx, Hadlima, or Humira) is required for diagnosis of hidradenitis suppurativa

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of uveitis

Biologic Immunomodulators PA - Yusimry

### Drug Name(s)

Yusimry

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

# Other Criteria:

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of juvenile idiopathic arthritis

Use of TWO preferred agents (Enbrel, Hadlima, or Humira) is required for diagnosis of rheumatoid arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, or Humira) is required for diagnosis of ankylosing spondylitis

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of psoriatic arthritis

Use of TWO preferred agents (Cosentyx, Enbrel, Hadlima, Humira, Otezla, Skyrizi, Stelara, or Tremfya) is required for diagnosis of plaque psoriasis

Use of TWO preferred agents (Hadlima, Humira, or Stelara) is required for diagnosis of ulcerative colitis

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of pediatric Crohn's disease

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Use of TWO preferred agents (Cosentyx, Hadlima, or Humira) is required for diagnosis of hidradenitis suppurativa

Use of TWO preferred agents (Hadlima and Humira) is required for diagnosis of uveitis

Biologic Immunomodulators PA - Zymfentra

### Drug Name(s)

Zymfentra 1-Pen

Zymfentra 2-Pen

Zymfentra 2-Syringe

### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. ONE of the following:
    - i. Patient's diagnosis is indicated for preferred biologic immunomodulator agent(s) AND ONE of the following:
      - a. Patient's medication history indicates use of preferred biologic immunomodulator agent(s) OR
      - b. Patient has an intolerance or hypersensitivity to preferred biologic immunomodulator agent(s) OR
      - c. Patient has an FDA labeled contraindication to preferred biologic immunomodulator agent(s) OR
    - ii. The request is for an FDA labeled indication that is not covered by preferred biologic immunomodulator agent(s) AND
- 3. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical improvement (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. Patient will NOT be using the requested agent in combination with another biologic immunomodulator AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

# **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

# Other Criteria:

Use of TWO preferred agents (Hadlima, Humira, Skyrizi, or Stelara) is required for diagnosis of adult Crohn's disease

Use of TWO preferred agents (Humira, Hadlima, or Stelara) is required for diagnosis of ulcerative colitis

NO preferred agent is required for diagnosis of adult fistulizing Crohn's disease

Bivigam/Flebogamma/Gammaplex/Octagam/Privigen PA

# Drug Name(s)

Bivigam

Flebogamma Dif

Gammaplex

Octagam

Privigen

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Budesonide Oral ER PA – Entocort

# Drug Name(s)

Budesonide Dr (Entocort)

# **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Budesonide Oral ER PA – Uceris

# Drug Name(s)

Budesonide Er (Uceris)

Uceris

#### Indications:

All Medically-Accepted Indications.

# **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Bydureon PA

# Drug Name(s)

Bydureon Bcise

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

Requested agent will be used for weight loss alone

### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of type 2 diabetes mellitus AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 180 days OR
  - C. ALL of the following:
    - i. ONE of the following:
      - 1. Patient's medication history includes use of a non glucagon-like peptide-1 (GLP-1) oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) within the past 90 days OR
      - 2. Patient had an ineffective treatment response to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 3. Patient has an intolerance or hypersensitivity to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 4. Patient has an FDA labeled contraindication to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent AND iii. Patient will NOT be using the requested agent in combination with another GLP-1 agonist agent, or an agent containing a GLP-1 agonist AND
    - iv. Patient will NOT be using the requested agent in combination with an agent containing a dipeptidyl peptidase-4 (DPP-4) inhibitor

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Byetta PA

# Drug Name(s)

Byetta

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

Requested agent will be used for weight loss alone

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of type 2 diabetes mellitus AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 180 days OR
  - C. ALL of the following:
    - i. ONE of the following:
      - 1. Patient's medication history includes use of a non glucagon-like peptide-1 (GLP-1) oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) within the past 90 days OR
      - 2. Patient had an ineffective treatment response to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 3. Patient has an intolerance or hypersensitivity to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 4. Patient has an FDA labeled contraindication to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent AND iii. Patient will NOT be using the requested agent in combination with another GLP-1 agonist agent, or an agent containing a GLP-1 agonist AND
    - iv. Patient will NOT be using the requested agent in combination with an agent containing a dipeptidyl peptidase-4 (DPP-4) inhibitor

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Bylvay PA

# Drug Name(s)

Bylvay

Bylvay (Pellets)

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. Patient has a diagnosis of progressive familial intrahepatic cholestasis (PFIC) AND
    - ii. The requested agent will be used to treat pruritus OR
  - B. BOTH of the following:
    - i. Patient has a diagnosis of Alagille Syndrome (ALGS) AND
    - ii. The requested agent will be used to treat cholestatic pruritus AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

Patient is within the FDA labeled age for the requested agent

# **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, hepatologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Camzyos PA

# Drug Name(s)

Camzyos

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of symptomatic New York Heart Association (NYHA) class II-III obstructive hypertrophic cardiomyopathy (HCM) AND
- 2. The requested agent will be used to improve functional capacity and symptoms

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cardiologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be for 12 months

Carglumic PA

# Drug Name(s)

Carbaglu

Carglumic Acid

#### **Indications:**

All FDA-Approved Indications, Some Medically-Accepted Indications.

#### **Off-Label Uses:**

For generic carglumic acid only - Acute hyperammonemia due to propionic acidemia (PA) or methylmalonic acidemia (MMA)

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of ONE of the following:
  - a. Acute hyperammonemia due to the deficiency of the hepatic enzyme N-acetylglutamate synthase (NAGS) OR
  - b. Chronic hyperammonemia due to the deficiency of the hepatic enzyme N-acetylglutamate synthase (NAGS) OR
  - c. Acute hyperammonemia due to propionic acidemia (PA) or methylmalonic acidemia (MMA) AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., geneticist, nephrologist, metabolic disorders) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Cayston PA

Drug Name(s)

Cayston

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Chenodal PA

Drug Name(s)

Chenodal

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of radiolucent stones in a well-opacifying gallbladder AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Chorionic Gonadotropin PA

### Drug Name(s)

Chorionic Gonadotropin

Novarel

Pregnyl

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

Requested agent will be used to promote fertility AND requested agent will be used to treat erectile dysfunction AND FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of prepubertal cryptorchidism not due to anatomic obstruction OR
  - B. Patient's sex is male, with a diagnosis of hypogonadotropic hypogonadism (hypogonadism secondary to pituitary deficiency) AND BOTH of the following:
    - i. Patient has a measured current or pretreatment total serum testosterone level that is below the testing laboratory's lower limit of the normal range or is less than 300  $\,$ ng/dL OR a free serum testosterone level that is below the testing laboratory's lower limit of the normal range AND
    - ii. Patient has measured luteinizing hormone (LH) AND follicle-stimulating hormone (FSH) levels that are at (low-normal) or below the testing laboratory's normal range OR
  - C. Patient has an indication that is supported in CMS approved compendia for the requested agent

**Age Restriction:** 

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Cibingo PA

# Drug Name(s)

Cibingo

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of moderate-to-severe atopic dermatitis AND BOTH of the following:
  - A. ONE of the following:
    - i. Patient has tried and failed a topical steroid (e.g., triamcinolone) OR
    - ii. Patient has an intolerance or hypersensitivity to a topical steroid OR
    - iii. Patient has an FDA labeled contraindication to a topical steroid AND
  - B. ONE of the following:
    - i. Patient has tried and failed a topical calcineurin inhibitor (e.g., tacrolimus) OR
    - ii. Patient has an intolerance or hypersensitivity to a topical calcineurin inhibitor OR
    - iii. Patient has an FDA labeled contraindication to a topical calcineurin inhibitor AND
- 2. Patient will NOT be using the requested agent in combination with other biologic immunomodulator agents, other JAK inhibitors, OR other immunosuppressants

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of moderate-to-severe atopic dermatitis AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with other biologic immunomodulator agents, other JAK inhibitors, OR other immunosuppressants

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, dermatologist, immunologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be for 12 months

Cinacalcet PA

### Drug Name(s)

Cinacalcet Hcl

Sensipar

### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

- 1. Patient has ONE of the following:
  - A. A diagnosis of hypercalcemia due to parathyroid carcinoma OR
  - B. A diagnosis of primary hyperparathyroidism (HPT) AND BOTH of the following:
    - i. Patient has a pretreatment serum calcium level that is above the testing laboratory's upper limit of normal AND
    - ii. Patient is unable to undergo parathyroidectomy OR
  - C. Another indication that is FDA approved or supported in CMS approved compendia for the requested agent not otherwise excluded from Part D [i.e., secondary hyperparathyroidism due to end-stage renal disease (ESRD) on dialysis]

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Cingair PA

# Drug Name(s)

Cingair

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of severe asthma with an eosinophilic phenotype AND
- 2. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LTRA, LAMA, theophylline) in combination with the requested agent AND
- 3. Patient will NOT be using the requested agent in combination with Xolair, Dupixent, or with another injectable interleukin 5 (IL-5) inhibitor (e.g., Fasenra, Nucala) for the requested indication AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of severe asthma with an eosinophilic phenotype AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LTRA, LAMA, theophylline) in combination with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with Xolair, Dupixent, or with another injectable interleukin 5 (IL-5) inhibitor (e.g., Fasenra, Nucala) for the requested indication AND
- 6. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

Patient is 18 years of age or over

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, immunologist, pulmonologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Colony Stimulating Factors PA – Fulphila

# Drug Name(s)

Fulphila

# **Indications:**

All Medically-Accepted Indications.

# **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

# **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Fylnetra

# Drug Name(s)

Fylnetra

# **Indications:**

All Medically-Accepted Indications.

# **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

# **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Granix

# Drug Name(s)

Granix

#### **Indications:**

All Medically-Accepted Indications.

# **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

# **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Leukine

# Drug Name(s)

Leukine

#### **Indications:**

All Medically-Accepted Indications.

# **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

# **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Neulasta

# Drug Name(s)

Neulasta

Neulasta Onpro Kit

#### **Indications:**

All Medically-Accepted Indications.

#### **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - i. Patient has an FDA labeled or CMS compendia supported indication for the requested agent that is not indicated for ONE preferred agent OR
  - ii. Patient has tried and had an inadequate response to ONE preferred agent OR
  - iii. Patient has an intolerance or hypersensitivity to ONE preferred agent OR
  - iv. Patient has an FDA labeled contraindication to ONE preferred agent OR
  - v. Prescriber has provided information in support of the use of the non-preferred agent over ONE preferred agent

Preferred agent(s) are Fulphila, Fylnetra, Granix, Nivestym, Nyvepria, Releuko, Stimufend, Udenyca, Zarxio, Ziextenzo

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Neupogen

### Drug Name(s)

Neupogen

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - i. Patient has an FDA labeled or CMS compendia supported indication for the requested agent that is not indicated for ONE preferred agent OR
  - ii. Patient has tried and had an inadequate response to ONE preferred agent OR
  - iii. Patient has an intolerance or hypersensitivity to ONE preferred agent OR
  - iv. Patient has an FDA labeled contraindication to ONE preferred agent OR
  - v. Prescriber has provided information in support of the use of the non-preferred agent over ONE preferred agent

Preferred agent(s) are Fulphila, Fylnetra, Granix, Nivestym, Nyvepria, Releuko, Stimufend, Udenyca, Zarxio, Ziextenzo

#### Age Restriction:

# **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Nivestym

# Drug Name(s)

Nivestym

### **Indications:**

All Medically-Accepted Indications.

# **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

# **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Nyvepria

# Drug Name(s)

Nyvepria

## **Indications:**

All Medically-Accepted Indications.

## **Off-Label Uses:**

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA - Releuko

# Drug Name(s)

Releuko

#### **Indications:**

All Medically-Accepted Indications.

## **Off-Label Uses:**

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

# **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Stimufend

# Drug Name(s)

Stimufend

#### Indications:

All Medically-Accepted Indications.

## **Off-Label Uses:**

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Udenyca

# Drug Name(s)

Udenyca

Udenyca Onbody

#### **Indications:**

All Medically-Accepted Indications.

# Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Zarxio

# Drug Name(s)

Zarxio

#### **Indications:**

All Medically-Accepted Indications.

## **Off-Label Uses:**

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Colony Stimulating Factors PA – Ziextenzo

# Drug Name(s)

Ziextenzo

#### **Indications:**

All Medically-Accepted Indications.

## **Off-Label Uses:**

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, hematologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 6 months

Concerta PA

# Drug Name(s)

Concerta

Methylphenidate Hcl Er (Concerta)

Indications:

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Corlanor PA

# Drug Name(s)

Corlanor

**Ivabradine** 

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has stable, symptomatic chronic heart failure (e.g., NYHA Class II, III, IV: ACCF/AHA Class C, D) AND
- 2. ONE of following:
  - A. ALL of the following:
    - i. The requested agent is for a pediatric patient, 6 months of age or over AND
    - ii. Patient has heart failure due to dilated cardiomyopathy (DCM) AND
    - iii. Patient is in sinus rhythm with an elevated heart rate OR
  - B. ALL of the following:
    - i. The requested agent is for an adult patient AND
    - ii. Patient has a baseline OR current left ventricular ejection fraction of 35% or less AND
    - iii. Patient is in sinus rhythm with a resting heart rate of 70 beats or greater per minute prior to initiating therapy with the requested agent AND
    - iv. ONE of the following:
      - a. Patient is on a maximally tolerated dose of beta blocker (e.g., bisoprolol, carvedilol, metoprolol) OR
      - b. Patient has an intolerance, FDA labeled contraindications, or hypersensitivity to a beta blocker

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Cortrophin Gel PA

Drug Name(s)

Cortrophin

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of nephrotic syndrome AND ONE of the following:
    - i. Patient has failed a conventional agent (i.e., prednisone, tacrolimus) for the requested indication OR
    - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to a conventional agent OR
  - B. Patient has a diagnosis of multiple sclerosis AND ALL of the following:
    - i. Patient is experiencing an acute exacerbation AND
    - ii. If indicated, there is evidence of a claim that the patient is currently being treated with a disease modifying drug (DMD) within the past 90 days [e.g., Avonex, dimethyl fumarate, glatiramer] to control disease progression OR has an intolerance, FDA labeled contraindication, or hypersensitivity to a DMD AND
    - iii. ONE of the following:
      - 1. Patient has failed corticosteroid therapy (e.g., methylprednisolone) within the last 30 days OR
      - 2. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to corticosteroid therapy OR

Criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 1 month

- C. Patient has a diagnosis of rheumatic disorder (e.g., ankylosing spondylitis, juvenile idiopathic arthritis, juvenile rheumatoid arthritis, psoriatic arthritis, rheumatoid arthritis) AND ALL of the following:
  - i. The requested agent will be used as adjunct therapy for short-term administration (to tide the patient over an acute episode or exacerbation) AND
  - ii. There is evidence of a claim that the patient is currently being treated with a conventional agent within the past 90 days [e.g., DMARD (methotrexate, leflunomide), biologics (Hadlima)] to control disease progression AND
  - iii. ONE of the following:

- 1. Patient has failed corticosteroid therapy (e.g., methylprednisolone) within the last 30 days OR
- 2. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to corticosteroid therapy OR
- D. Patient has a diagnosis of systemic lupus erythematosus (SLE) disease AND the patient will continue standard SLE therapy [corticosteroids (e.g., methylprednisolone, prednisone), hydroxychloroquine, immunosuppressives (e.g., azathioprine, methotrexate, oral cyclophosphamide)] in combination with the requested agent OR
- E. Patient has another FDA approved indication AND ONE of the following:
  - i. Patient has failed corticosteroid therapy (e.g., methylprednisolone) within the last 30 days OR
  - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to corticosteroid therapy OR
- F. Patient has another indication that is supported in CMS approved compendia for the requested agent AND ONE of the following:
  - i. Patient has failed corticosteroid therapy (e.g., methylprednisolone) within the last 30 days OR
  - ii. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to corticosteroid therapy AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Cotempla PA

Drug Name(s)

Cotempla Xr-Odt

**Indications:** 

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Cresemba PA

# Drug Name(s)

Cresemba

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of invasive aspergillosis OR
  - B. Patient has a diagnosis of invasive mucormycosis OR
  - C. Patient has another indication that is supported in CMS approved compendia for the requested agent

Criteria for renewal approval require BOTH of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of invasive aspergillosis and patient has continued indicators of active disease (e.g., continued radiologic findings, direct microscopy findings, histopathology findings, positive cultures, positive serum galactomannan assay) OR
  - B. Patient has a diagnosis of invasive mucormycosis and patient has continued indicators of active disease (e.g., continued radiologic findings, direct microscopy findings, histopathology findings, positive cultures, positive serum galactomannan assay) OR
  - C. BOTH of the following:
    - i. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient has had clinical benefit with the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 6 months

Crinone PA

# Drug Name(s)

Crinone

#### Indications:

All Medically-Accepted Indications.

## **Off-Label Uses:**

#### **Exclusion Criteria:**

Requested agent will be used to treat infertility AND FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Crysvita PA

# Drug Name(s)

Crysvita

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. Patient has a diagnosis of X-linked hypophosphatemia (XLH) as confirmed by ONE of the following:
      - a. Genetic testing OR
      - b. Elevated levels of intact fibroblast growth factor 23 (FGF23) OR
      - c. Prescriber has provided information indicating the patient has a positive family history of XLH AND
    - ii. ONE of the following:
      - a. Patient's epiphyseal plate has not fused OR
      - b. Patient's epiphyseal plate has fused AND the patient is experiencing symptoms of XLH (e.g., bone pain, fractures, limited mobility) OR
  - B. Patient has a diagnosis of tumor-induced osteomalacia (TIO) associated with phosphaturic mesenchymal tumors AND BOTH of the following:
    - i. The requested agent is being used to treat FGF23 related hypophosphatemia AND
    - ii. The tumor cannot be curatively surgically resected or localized AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of X-linked hypophosphatemia (XLH) OR
  - B. Patient has a diagnosis of tumor-induced osteomalacia (TIO) associated with phosphaturic mesenchymal tumors AND
- 3. Patient has had clinical benefit with the requested agent (e.g., enhanced height velocity, improvement in lower extremity bowing and associated abnormalities, radiographic evidence of epiphyseal healing, improvement in bone pain, enhanced mobility, improvement in osteomalacia, improvement in fracture healing) AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

Patient is within the FDA labeled age for the requested agent for the requested indication **Prescriber Restrictions:** 

Prescriber is a specialist in the area of the patient's diagnosis (e.g., nephrologist, endocrinologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

**Coverage Duration:** 

Approval will be for 12 months

Cutaquig PA

Drug Name(s)

Cutaquig

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require ONE of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Primary immunodeficiency [e.g., congenital agammaglobulinemia, common variable immunodeficiency (CVID), severe combined immunodeficiency, Wiskott-Aldrich Syndrome, X-linked agammaglobulinemia (XLA), humoral immunodeficiency, IgG subclass deficiency with or without IgA deficiency] OR
  - B. Multiple sclerosis (MS) AND BOTH of the following:
    - i. Patient has a diagnosis of relapsing remitting MS (RRMS) AND
    - ii. Patient has had an insufficient response, documented failure, or FDA labeled contraindication to TWO MS agents (e.g., Avonex, Betaseron, Copaxone, dimethyl fumarate, fingolimod, glatiramer, Glatopa, Kesimpta, Plegridy, teriflunomide, Vumerity) OR
- 2. ONE of the following:
  - A. Patient has another FDA labeled indication for the requested agent OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

Drug is also subject to Part B versus Part D review.

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

**Cuvrior PA** 

## Drug Name(s)

Cuvrior

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of Wilson's disease confirmed by ONE of the following:
  - A. Confirmation of genetic mutation of the ATP7B gene OR
  - B. Patient has TWO of the following:
    - i. Presence of hepatic abnormality (e.g., acute liver failure, cirrhosis, fatty liver)
    - ii. Presence of Kayser-Fleischer rings
    - iii. Serum ceruloplasmin level less than 20 mg/dL
    - iv. Basal urinary copper excretion greater than 40 mcg/24 hours or the testing laboratory's upper limit of normal
    - v. Hepatic parenchymal copper content greater than 40 mcg/g dry weight
    - vi. Presence of neurological symptoms (e.g., dystonia, hypertonia, rigidity with tremors, dysarthria, muscle spasms, dysphasia, polyneuropathy, dysautonomia) AND
- 2. Patient is decoppered AND
- 3. Patient has stable Wilson's disease AND
- 4. Patient is tolerant of penicillamine

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Wilson's disease AND
- 3. Patient has had clinical benefit with the requested agent

### Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, hepatologist, neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Cystadrops PA

Drug Name(s)

Cystadrops

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., ophthalmologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Cystaran PA

# Drug Name(s)

Cystaran

# Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., ophthalmologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Cystinosis Agents PA – Cystagon

# Drug Name(s)

Cystagon

#### **Indications:**

All FDA-Approved Indications.

#### **Off-Label Uses:**

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of nephropathic cystinosis AND
- 2. Prescriber has performed a baseline white blood cell (WBC) cystine level test AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of nephropathic cystinosis AND
- 3. Patient has had clinical benefit with the requested agent (e.g., decrease in WBC cystine levels from baseline) AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

Cystinosis Agents PA - Procysbi

# Drug Name(s)

Procysbi

#### **Indications:**

All FDA-Approved Indications.

#### **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of nephropathic cystinosis AND
- 2. Prescriber has performed a baseline white blood cell (WBC) cystine level test AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of nephropathic cystinosis AND
- 3. Patient has had clinical benefit with the requested agent (e.g., decrease in WBC cystine levels from baseline) AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

Dalfampridine PA

### Drug Name(s)

Ampyra

Dalfampridine Er

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of multiple sclerosis (MS) AND
- 2. ONE of the following:
  - A. The requested agent will be used in combination with a disease modifying agent [e.g., dimethyl fumarate, glatiramer (e.g., Copaxone)] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to a disease modifying agent OR
  - C. Prescriber has provided information indicating that a disease modifying agent is not clinically appropriate for the patient

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of multiple sclerosis (MS) AND
- 3. ONE of the following:
  - A. The requested agent will be used in combination with a disease modifying agent [e.g., dimethyl fumarate, glatiramer (e.g., Copaxone)] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to a disease modifying agent OR
  - C. Prescriber has provided information indicating that a disease modifying agent is not clinically appropriate for the patient AND
- 4. Patient has had improvements or stabilization from baseline in timed walking speed (timed 25-foot walk)

#### Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Initial approval will be for 3 months, renewal approval will be for 12 months

Daybue PA

## Drug Name(s)

Daybue

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require the following:

1. Patient has a diagnosis of Rett syndrome (RTT) with genetic analysis confirming mutation in the MECP2 gene

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Rett syndrome (RTT) AND
- 3. Patient has had clinical benefit with the requested agent

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., geneticist, neurologist, pediatrician) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Dayvigo PA

Drug Name(s)

Dayvigo

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Deferiprone PA

Drug Name(s)

Deferiprone

Ferriprox

Ferriprox Twice-A-Day

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of transfusional iron overload due to thalassemia syndromes, sickle cell disease or other anemias AND
- 2. Patient will NOT be using the requested agent in combination with another iron chelating agent (e.g., deferasirox) for the requested indication

# Age Restriction:

Patient is within the FDA labeled age for the requested agent for the requested indication

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Dichlorphenamide PA

# Drug Name(s)

Dichlorphenamide

Keveyis

Ormalvi

## **Indications:**

All FDA-Approved Indications.

## **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require the following:

1. Patient has a diagnosis of primary hyperkalemic periodic paralysis, primary hypokalemic periodic paralysis, or a related variant

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Doptelet PA

## Drug Name(s)

Doptelet

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of thrombocytopenia AND ALL of the following:
    - i. Patient has chronic liver disease AND
    - ii. Patient has a platelet count less than 50 X 10^9/L AND
    - iii. Patient is scheduled to undergo a procedure with an associated risk of bleeding (e.g., gastrointestinal endoscopy, liver biopsy, bronchoscopy, dental procedure) AND
    - iv. The length of therapy of the requested agent is within the FDA labeled duration for the requested indication OR
  - B. Patient has a diagnosis of chronic immune (idiopathic) thrombocytopenia (ITP) AND ONE of the following:
    - i. Patient has tried and had an insufficient response to a corticosteroid, another thrombopoietin receptor agonist (e.g., Promacta), or immunoglobulin (IVIg or anti-D) OR ii. Patient has an intolerance or hypersensitivity to a corticosteroid, another thrombopoietin receptor agonist (e.g., Promacta), or immunoglobulin (IVIg or anti-D) OR iii. Patient has an FDA labeled contraindication to a corticosteroid, another thrombopoietin receptor agonist (e.g., Promacta), or immunoglobulin (IVIg or anti-D) OR
- iv. Patient has had an insufficient response to a splenectomy AND 2. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Initial (I) 6 mo ITP, Renewal (R) 12 mo ITP. I & R 1 mo thrombocytopenia w/chronic liver disease

#### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of thrombocytopenia AND ALL of the following:
    - i. Patient has chronic liver disease AND
    - ii. Patient has a platelet count less than 50 X 10^9/L AND
    - iii. Patient is scheduled to undergo a procedure with an associated risk of bleeding (e.g., gastrointestinal endoscopy, liver biopsy, bronchoscopy, dental procedure) AND
    - iv. The length of therapy of the requested agent is within the FDA labeled duration for the requested indication OR

- B. Patient has a diagnosis of chronic immune (idiopathic) thrombocytopenia (ITP) AND ONE of the following:
  - i. Patient's platelet count is 50 x 10^9/L or greater OR
  - ii. Patient's platelet count has increased sufficiently to avoid clinically significant bleeding AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Droxidopa PA

## Drug Name(s)

Droxidopa

Northera

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of neurogenic orthostatic hypotension (nOH) AND
- 2. Prescriber has performed baseline blood pressure readings while the patient is sitting or supine (lying face up), AND also within three minutes of standing from a supine position AND
- 3. Patient has a decrease of at least 20 mmHg in systolic blood pressure or 10 mmHg diastolic blood pressure within three minutes after standing AND
- 4. Patient has persistent and consistent symptoms of neurogenic orthostatic hypotension (nOH) caused by ONE of the following:
  - A. Primary autonomic failure [Parkinson's disease (PD), multiple system atrophy, or pure autonomic failure] OR
  - B. Dopamine beta-hydroxylase deficiency OR
  - C. Non-diabetic autonomic neuropathy AND
- 5. Prescriber has assessed the severity of the patient's baseline symptoms of dizziness, lightheadedness, feeling faint, or feeling like the patient may black out AND
- 6. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of neurogenic orthostatic hypotension (nOH) AND
- 3. Patient has had improvements or stabilization with the requested agent as indicated by improvement in severity from baseline symptoms of ONE of the following:
  - A. Dizziness
  - B. Lightheadedness
  - C. Feeling faint
  - D. Feeling like the patient may black out AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cardiologist, neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be 1 month for initial, 3 months for renewal

**Dupixent PA** 

## Drug Name(s)

Dupixent

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of moderate-to-severe atopic dermatitis AND ALL of the following:
    - i. ONE of the following:
      - a. Patient has tried and failed a topical steroid (e.g., triamcinolone) OR
      - b. Patient has an intolerance, hypersensitivity, or an FDA labeled contraindication to a topical steroid AND
    - ii. For patients 2 years of age or over, ONE of the following:
      - a. Patient has tried and failed a topical calcineurin inhibitor (e.g., tacrolimus) OR
      - b. Patient has an intolerance, hypersensitivity, or an FDA labeled contraindication to a topical calcineurin inhibitor AND
    - iii. Patient will NOT be using the requested agent in combination with another biologic agent or a JAK inhibitor for the requested indication OR
  - B. Patient has a diagnosis of moderate-to-severe asthma with an eosinophilic phenotype or oral corticosteroid dependent asthma AND BOTH of the following:
    - i. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LRTA, LAMA, theophylline) in combination with the requested agent AND
    - ii. Patient will NOT be using the requested agent in combination with Xolair or with an injectable interleukin 5 (IL-5) inhibitor (e.g., Cinqair, Fasenra, Nucala) for the requested indication OR
  - C. Patient has a diagnosis of chronic rhinosinusitis with nasal polyposis (CRSwNP) AND the following:
    - i. BOTH of the following:
      - a. ONE of the following:
        - 1. Patient has tried and had an inadequate response to an oral systemic corticosteroid AND an intranasal corticosteroid (e.g., fluticasone) OR
        - 2. Patient has an intolerance, hypersensitivity, or an FDA labeled contraindication to an oral systemic corticosteroid AND an intranasal corticosteroid AND

Initial criteria continues: see Other Criteria

# Age Restriction:

For diagnosis of moderate-to-severe atopic dermatitis, patient is 6 months of age or over. For diagnosis of moderate-to-severe asthma with an eosinophilic phenotype or oral corticosteroid dependent asthma,

patient is 6 years of age or over. For diagnosis of CRSwNP, patient is 18 years of age or over. For diagnosis of EoE, patient is 1 year of age or over. For diagnosis of PN, patient is 18 years of age or over.

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, dermatologist, immunologist, gastroenterologist, otolaryngologist, pulmonologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

- b. Patient will continue standard maintenance therapy (e.g., intranasal corticosteroid) in combination with the requested agent OR
- D. Patient has a diagnosis of eosinophilic esophagitis (EoE) confirmed by esophageal biopsy OR
- E. Patient has a diagnosis of prurigo nodularis (PN) AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of moderate-to-severe atopic dermatitis AND the following:
    - i. Patient will NOT be using the requested agent in combination with another biologic agent or a JAK inhibitor for the requested indication OR
  - B. Patient has a diagnosis of moderate-to-severe asthma with an eosinophilic phenotype or oral corticosteroid dependent asthma AND BOTH of the following:
    - i. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LTRA, LAMA, theophylline) in combination with the requested agent AND ii. Patient will NOT be using the requested agent in combination with Xolair or with an injectable interleukin 5 (IL-5) inhibitor (e.g., Cinqair, Fasenra, Nucala) for the requested indication OR
  - C. Patient has a diagnosis of chronic rhinosinusitis with nasal polyposis (CRSwNP) AND the following:
    - i. Patient will continue standard maintenance therapy (e.g., intranasal corticosteroid) in combination with the requested agent OR
  - D. Patient has a diagnosis of eosinophilic esophagitis (EoE) OR
  - E. Patient has a diagnosis of prurigo nodularis (PN) AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Elmiron PA

# Drug Name(s)

Elmiron

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. The requested agent will be used for the relief of bladder pain or discomfort associated with interstitial cystitis AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. The requested agent will be used for the relief of bladder pain or discomfort associated with interstitial cystitis AND
- 3. Patient has had clinical benefit with the requested agent (e.g., decreased bladder pain, decreased frequency or urgency of urination) AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be 6 months for initial, 12 months for renewal

Emflaza PA

### Drug Name(s)

Deflazacort

**Emflaza** 

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of Duchenne muscular dystrophy (DMD) confirmed by ONE of the following:
  - A. Presence of abnormal dystrophin OR
  - B. Confirmed mutation of the dystrophin gene AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Duchenne muscular dystrophy (DMD) AND
- 3. Patient has had improvement, stabilization of the disease, or clinical benefit from baseline (e.g., improved strength and timed motor function, improved pulmonary function, reduced the need for scoliosis surgery) AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

**Emgality PA** 

## Drug Name(s)

Emgality

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of migraine AND ALL of the following:
    - i. The requested agent is being used for migraine prophylaxis AND
    - ii. Patient has 4 or more migraine headache days per month AND
    - iii. Patient will NOT be using the requested agent in combination with another calcitonin gene-related peptide (CGRP) agent for migraine prophylaxis OR
  - B. Patient has a diagnosis of episodic cluster headache AND BOTH of the following:
    - i. Patient has had at least 5 cluster headache attacks AND
    - ii. Patient has had at least two cluster periods lasting 7 days to one year and separated by pain-free remission periods of 3 months or more

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. ALL of the following:
    - i. Patient has a diagnosis of migraine AND
    - ii. The requested agent is being used for migraine prophylaxis AND
    - iii. Patient will NOT be using the requested agent in combination with another calcitonin gene-related peptide (CGRP) agent for migraine prophylaxis OR
  - B. Patient has a diagnosis of episodic cluster headache AND
- 3. Patient has had clinical benefit with the requested agent

**Age Restriction:** 

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Emsam PA

### Drug Name(s)

Emsam

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of major depressive disorder (MDD) OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. BOTH of the following:
    - i. ONE of the following:
      - a. BOTH of the following:
        - i. Patient has a diagnosis of major depressive disorder (MDD) AND
        - ii. ONE of the following:
          - 1. Patient has tried and had an inadequate response to at least two different oral antidepressants (e.g., SSRIs, SNRIs, mirtazapine, bupropion) OR
          - 2. Patient has an intolerance or hypersensitivity to at least two different oral antidepressants (e.g., SSRIs, SNRIs, mirtazapine, bupropion) OR
          - 3. Patient has an FDA labeled contraindication to at least two different oral antidepressants (e.g., SSRIs, SNRIs, mirtazapine, bupropion) OR
      - b. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent

# Age Restriction:

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:

- A. Patient has a diagnosis of major depressive disorder (MDD) OR
- B. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
- 3. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. BOTH of the following:
    - i. Patient has had clinical benefit with the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent

Endari PA

### Drug Name(s)

Endari

L-Glutamine

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of sickle cell disease AND
- 2. Patient is using the requested agent to reduce the acute complications of sickle cell disease AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to maximally tolerated dose of hydroxyurea OR
  - B. Patient has an intolerance or hypersensitivity to hydroxyurea OR
  - C. Patient has an FDA labeled contraindication to hydroxyurea AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of sickle cell disease AND
- 3. Patient is using the requested agent to reduce the acute complications of sickle cell disease AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

## **Prescriber Restrictions:**

### **Coverage Duration:**

Approval will be for 12 months

**Enspryng PA** 

## Drug Name(s)

Enspryng

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of neuromyelitis optica spectrum disorder (NMOSD) AND
- 2. Patient is anti-aquaporin-4 (AQP4) antibody positive AND
- 3. Prescriber has screened the patient for hepatitis B viral (HBV) infection and determined to NOT have active hepatitis B viral infection AND
- 4. Patient does NOT have active or untreated tuberculosis AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of neuromyelitis optica spectrum disorder (NMOSD) AND
- 3. Patient has had clinical benefit with the requested agent (e.g., decreased relapses, improvement or stabilization of vision or paralysis) AND
- 4. Patient does NOT have active hepatitis B virus (HBV) infection AND
- 5. Patient does NOT have active or untreated tuberculosis AND
- 6. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Eohilia PA

## Drug Name(s)

Eohilia

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of eosinophilic esophagitis (EoE) confirmed by esophageal biopsy AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, gastroenterologist, immunologist, pulmonologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 3 months

Epclusa PA

## Drug Name(s)

**Epclusa** 

Sofosbuvir/Velpatasvir

#### **Indications:**

All Medically-Accepted Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of hepatitis C confirmed by serological markers OR
  - B. Patient is a hepatitis C virus (HCV) uninfected solid organ transplant recipient AND BOTH of the following:
    - i. Patient received an HCV viremic donor organ AND
    - ii. The requested agent is being used for prophylaxis AND
- 2. Prescriber has screened the patient for current or prior hepatitis B viral (HBV) infection and if positive, will monitor the patient for HBV flare-up or reactivation during and after treatment with the requested agent AND
- 3. The requested agent will be used in a treatment regimen and length of therapy that is supported in FDA approved labeling or AASLD/IDSA guidelines for the patient's diagnosis and genotype AND
- 4. The requested dose is within FDA labeled dosing or supported in AASLD/IDSA guideline dosing for the requested indication AND
- 5. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
  - C. Patient has an FDA labeled contraindication or hypersensitivity to the preferred agent: Mavyret for supported genotypes OR
  - D. Prescriber has provided information based on FDA approved labeling or AASLD/IDSA guidelines supporting the use of the non-preferred agent for the patient's diagnosis and genotype over the preferred agent: Mavyret for supported genotypes

### **Age Restriction:**

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, hepatologist or infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Duration of therapy: Based on FDA approved labeling or AASLD/IDSA guideline supported **Other Criteria**:

**Epidiolex PA** 

## Drug Name(s)

Epidiolex

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of seizures associated with ONE of the following:
  - A. Lennox-Gastaut syndrome OR
  - B. Dravet syndrome OR
  - C. Tuberous sclerosis complex AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Erythropoietin Stimulating Agents PA – Aranesp

## Drug Name(s)

Aranesp Albumin Free

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. The requested agent is being prescribed for ONE of the following:
  - A. Anemia due to myelosuppressive chemotherapy for a non-myeloid malignancy AND ALL of the following:
    - i. Patient's hemoglobin level is less than 10 g/dL for patients initiating ESA therapy OR less than 12 g/dL for patients stabilized on therapy (measured within the previous 4 weeks) AND
    - ii. Patient is being concurrently treated with chemotherapy with or without radiation (there must be a minimum of 2 additional months of planned chemotherapy) AND
    - iii. The intent of chemotherapy is non-curative OR
  - B. Anemia associated with chronic kidney disease in a patient NOT on dialysis AND ALL of the following:
    - i. Patient's hemoglobin level is less than 10 g/dL for patients initiating ESA therapy OR 11 g/dL or less for patients stabilized on therapy (measured within the previous 4 weeks) AND
    - ii. The rate of hemoglobin decline indicates the likelihood of requiring a RBC transfusion AND
    - iii. The intent of therapy is to reduce the risk of alloimmunization and/or other RBC transfusion related risks OR
  - C. Anemia due to myelodysplastic syndrome AND the patient's hemoglobin level is less than 12 g/dL for patients initiating ESA therapy OR less than or equal to 12 g/dL for patients stabilized on therapy (measured within the previous 4 weeks) OR
  - D. Another indication that is supported in CMS approved compendia for the requested agent AND the patient's hemoglobin level is less than 12 g/dL for patients initiating ESA therapy OR less than or equal to 12 g/dL for patients stabilized on therapy (measured within the previous 4 weeks) AND
- 2. Patient's transferrin saturation and serum ferritin have been evaluated

Drug is also subject to Part B versus Part D review.

#### Age Restriction:

## **Prescriber Restrictions:**

#### **Coverage Duration:**

6 months for chemotherapy, 12 months for other indications

Erythropoietin Stimulating Agents PA - Epogen/Procrit

### Drug Name(s)

Epogen

Procrit

#### **Indications:**

All Medically-Accepted Indications.

#### **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. The requested agent is being prescribed for ONE of the following:
  - A. To reduce the possibility of allogeneic blood transfusion in a surgery patient AND the patient's hemoglobin level is greater than 10 g/dL but less than or equal to 13 g/dL OR
  - B. Anemia due to myelosuppressive chemotherapy for a non-myeloid malignancy AND ALL of the following:
    - i. Patient's hemoglobin level is less than 10 g/dL for patients initiating ESA therapy OR less than 12 g/dL for patients stabilized on therapy (measured within the previous 4 weeks) AND
    - ii. Patient is being concurrently treated with chemotherapy with or without radiation (there must be a minimum of 2 additional months of planned chemotherapy) AND
    - iii. The intent of chemotherapy is non-curative OR
  - C. Anemia associated with chronic kidney disease in a patient NOT on dialysis AND ALL of the following:
    - i. Patient's hemoglobin level is less than 10 g/dL for patients initiating ESA therapy OR 11 g/dL or less for patients stabilized on therapy (measured within the previous 4 weeks) AND
    - ii. The rate of hemoglobin decline indicates the likelihood of requiring a RBC transfusion AND
    - iii. The intent of therapy is to reduce the risk of alloimmunization and/or other RBC transfusion related risks OR
  - D. Anemia due to myelodysplastic syndrome AND the patient's hemoglobin level is less than 12 g/dL for patients initiating ESA therapy OR less than or equal to 12 g/dL for patients stabilized on therapy (measured within the previous 4 weeks) OR
  - E. Anemia resulting from zidovudine treatment of HIV infection AND the patient's hemoglobin level is less than 12 g/dL for patients initiating ESA therapy OR less than or equal to 12 g/dL for patients stabilized on therapy (measured within the previous 4 weeks) OR

Initial criteria continues: see Other Criteria

Age Restriction:

Prescriber Restrictions: Coverage Duration:

1 month for surgery (reduce transfusion possibility), 6 months for chemo, 12 months for other **Other Criteria:** 

F. Another indication that is supported in CMS approved compendia for the requested agent AND the patient's hemoglobin level is less than 12 g/dL for patients initiating ESA therapy OR less than or equal to 12 g/dL for patients stabilized on therapy (measured within the previous 4 weeks) AND

2. Patient's transferrin saturation and serum ferritin have been evaluated

Drug is also subject to Part B versus Part D review.

Erythropoietin Stimulating Agents PA – Retacrit

## Drug Name(s)

Retacrit

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. The requested agent is being prescribed for ONE of the following:
  - A. To reduce the possibility of allogeneic blood transfusion in a surgery patient AND the patient's hemoglobin level is greater than 10 g/dL but less than or equal to 13 g/dL OR B. Anemia due to myelosuppressive chemotherapy for a non-myeloid malignancy AND ALL or
  - B. Anemia due to myelosuppressive chemotherapy for a non-myeloid malignancy AND ALL of the following:
    - i. Patient's hemoglobin level is less than 10 g/dL for patients initiating ESA therapy OR less than 12 g/dL for patients stabilized on therapy (measured within the previous 4 weeks) AND
    - ii. Patient is being concurrently treated with chemotherapy with or without radiation (there must be a minimum of 2 additional months of planned chemotherapy) AND iii. The intent of chemotherapy is non-curative OR
  - C. Anemia associated with chronic kidney disease in a patient NOT on dialysis AND ALL of the following:
    - i. Patient's hemoglobin level is less than 10 g/dL for patients initiating ESA therapy OR 11 g/dL or less for patients stabilized on therapy (measured within the previous 4 weeks) AND
    - ii. The rate of hemoglobin decline indicates the likelihood of requiring a RBC transfusion AND
    - iii. The intent of therapy is to reduce the risk of alloimmunization and/or other RBC transfusion related risks OR
  - D. Anemia resulting from zidovudine treatment of HIV infection AND the patient's hemoglobin level is less than 12 g/dL for patients initiating ESA therapy OR less than or equal to 12 g/dL for patients stabilized on therapy (measured within the previous 4 weeks) OR
  - E. Another indication that is supported in CMS approved compendia for the requested agent AND the patient's hemoglobin level is less than 12 g/dL for patients initiating ESA therapy OR less than or equal to 12 g/dL for patients stabilized on therapy (measured within the previous 4 weeks) AND
- 2. Patient's transferrin saturation and serum ferritin have been evaluated

Drug is also subject to Part B versus Part D review.

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

1 month for surgery (reduce transfusion possibility), 6 months for chemo, 12 months for other Other Criteria:

**Evenity PA** 

### Drug Name(s)

Evenity

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient is postmenopausal with a diagnosis of osteoporosis AND BOTH of the following:
  - A. Patient's diagnosis was confirmed by ONE of the following:
    - i. A fragility fracture in the hip or spine OR
    - ii. A T-score of -2.5 or lower OR
    - iii. A T-score of -1.0 to -2.5 AND ONE of the following:
      - a. A fragility fracture of the proximal humerus, pelvis, or distal forearm OR
      - b. A FRAX 10-year probability for major osteoporotic fracture of 20% or greater OR
      - c. A FRAX 10-year probability of hip fracture of 3% or greater AND
  - B. ONE of the following:
    - i. Patient is at a very high fracture risk as defined by ONE of the following:
      - a. Patient had a recent fracture (within the past 12 months) OR
      - b. Patient had fractures while on FDA approved osteoporosis therapy OR
      - c. Patient has had multiple fractures OR
      - d. Patient had fractures while on drugs causing skeletal harm (e.g., long-term glucocorticoids) OR
      - e. Patient has a very low T-score (less than -3.0) OR
      - f. Patient is at high risk for falls or has a history of injurious falls OR
      - g. Patient has a very high fracture probability by FRAX (e.g., major osteoporosis fracture greater than 30%, hip fracture greater than 4.5%) or by other validated fracture risk algorithm OR
    - ii. ONE of the following:
      - a. Patient has tried and had an inadequate response to a bisphosphonate OR
      - b. Patient has an intolerance or hypersensitivity to a bisphosphonate OR
      - c. Patient has an FDA labeled contraindication to a bisphosphonate AND

Criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

No prior use approve 12 months, Prior use approve remainder of 12 months of total cumulative therapy **Other Criteria:** 

2. ONE of the following:

- A. Patient has a pretreatment or current calcium level that is NOT below the lower limit of the testing laboratory's normal range OR
- B. Patient has a pretreatment or current calcium level that is below the lower limit of the testing laboratory's normal range AND it will be corrected prior to use of the requested agent OR
- C. Prescriber has indicated that the patient is not at risk for hypocalcemia (not including risk associated with the requested agent) AND
- 3. Patient will NOT be using the requested agent in combination with a bisphosphonate, denosumab (e.g., Prolia, Xgeva), or parathyroid hormone analog (e.g., abaloparatide, teriparatide) for the requested indication AND
- 4. The requested dose is within FDA labeled dosing for the requested indication AND
- 5. The total cumulative duration of treatment with Evenity (romosozumab-aqqg) has not exceeded 12 months

Evkeeza PA

### Drug Name(s)

Evkeeza

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of homozygous familial hypercholesterolemia (HoFH) AND
- 2. Patient's diagnosis was confirmed by ONE of the following:
  - A. Genetic confirmation of bi-allelic pathogenic/likely pathogenic variants on different chromosomes at the LDLR, Apo-B, PCSK9, or LDLRAP1 genes or greater than or equal to 2 such variants at different loci OR
  - B. History of untreated LDL-C greater than 400 mg/dL (greater than 10 mmol/L) AND ONE of the following:
    - i. Patient has cutaneous or tendon xanthomas before the age of 10 years OR
    - ii. Untreated elevated LDL-C levels consistent with heterozygous familial hypercholesterolemia (HeFH) in both parents (or in digenic form, one parent may have normal LDL-C levels and the other may have LDL-C levels consistent with HoFH) AND
- 3. ONE of the following:
  - A. Patient is currently being treated with a lipid-lowering regimen in the past 90 days (i.e., rosuvastatin in combination with ezetimibe, atorvastatin in combination with ezetimibe, OR PCSK9) OR
  - B. Patient has an intolerance or hypersensitivity to a lipid-lowering regimen (i.e., rosuvastatin in combination with ezetimibe, atorvastatin in combination with ezetimibe, OR PCSK9) OR
  - C. Patient has an FDA labeled contraindication to a lipid-lowering regimen (i.e., rosuvastatin in combination with ezetimibe, atorvastatin in combination with ezetimibe, OR PCSK9) AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

Patient is within FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cardiologist, endocrinologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of homozygous familial hypercholesterolemia (HoFH) AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. ONE of the following:

- A. Patient is currently being treated with a lipid-lowering regimen in the past 90 days (i.e., rosuvastatin in combination with ezetimibe, atorvastatin in combination with ezetimibe, OR PCSK9) OR
- B. Patient has an intolerance or hypersensitivity to a lipid-lowering regimen (i.e., rosuvastatin in combination with ezetimibe, atorvastatin in combination with ezetimibe, OR PCSK9)

  OR
- C. Patient has an FDA labeled contraindication to a lipid-lowering regimen (i.e., rosuvastatin in combination with ezetimibe, atorvastatin in combination with ezetimibe, OR PCSK9)

## AND

5. The requested dose is within FDA labeled dosing for the requested indication

Evrysdi PA

## Drug Name(s)

Evrysdi

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require the following:

1. Patient has a diagnosis of Spinal Muscular Atrophy (SMA) as confirmed by genetic testing

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Spinal Muscular Atrophy (SMA) AND
- 3. Patient has had clinical benefit with the requested agent

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist, geneticist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Eysuvis PA

## Drug Name(s)

Eysuvis

## Indications:

All FDA-Approved Indications.

### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of dry eye disease AND
- 2. The requested agent will be used for short-term (up to two weeks) treatment AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

**Prescriber Restrictions:** 

## **Coverage Duration:**

Approval will be for 1 month

Fabhalta PA

Drug Name(s)

Fabhalta

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Pending CMS Review

Fasenra PA

## Drug Name(s)

Fasenra

Fasenra Pen

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of severe asthma with an eosinophilic phenotype AND
- 2. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LTRA, LAMA, theophylline) in combination with the requested agent AND
- 3. Patient will NOT be using the requested agent in combination with Xolair, Dupixent, or with another injectable interleukin 5 (IL-5) inhibitor (e.g., Cinqair, Nucala) for the requested indication AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of severe asthma with an eosinophilic phenotype AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LTRA, LAMA, theophylline) in combination with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with Xolair, Dupixent, or with another injectable interleukin 5 (IL-5) inhibitor (e.g., Cinqair, Nucala) for the requested indication AND
- 6. The requested dose is within the FDA labeled dosing for the requested indication

#### Age Restriction:

Patient is 6 years of age or over

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, immunologist, pulmonologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Fentanyl Oral PA - Fentanyl lozenge

### Drug Name(s)

Fentanyl Citrate Oral Transmucosal

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. ONE of the following:
  - a. Patient has a documented diagnosis (i.e., medical records) of chronic cancer pain due to an active malignancy AND BOTH of the following:
    - i. Prescriber has provided the patient's type of cancer AND
    - ii. There is evidence of a claim that the patient is currently being treated with a longacting opioid with the requested agent within the past 90 days OR
  - b. Patient has a diagnosis that is supported in CMS approved compendia for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with any other oral or nasal fentanyl agent

## Age Restriction:

Patient is 16 years of age or over

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Fentanyl Oral PA - Fentora

### Drug Name(s)

Fentanyl Citrate Tablet

Fentora

#### **Indications:**

All Medically-Accepted Indications.

#### **Off-Label Uses:**

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. ONE of the following:
  - a. Patient has a documented diagnosis (i.e., medical records) of chronic cancer pain due to an active malignancy AND BOTH of the following:
    - i. Prescriber has provided the patient's type of cancer AND
    - ii. There is evidence of a claim that the patient is currently being treated with a longacting opioid with the requested agent within the past 90 days OR
  - b. Patient has a diagnosis that is supported in CMS approved compendia for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with any other oral or nasal fentanyl agent

## Age Restriction:

Patient is 18 years of age or over

## **Prescriber Restrictions:**

## **Coverage Duration:**

Approval will be for 12 months

Filspari PA

## Drug Name(s)

Filspari

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of primary immunoglobulin A nephropathy (IgAN) confirmed by kidney biopsy AND
- 2. The requested agent will be used to reduce proteinuria AND
- 3. Patient is at risk of rapid disease progression as shown by ONE of the following:
  - A. A urine protein-to-creatinine ratio (UPCR) greater than or equal to 0.8 g/g OR
  - B. Proteinuria greater than or equal to 1 g/day AND
- 4. ONE of the following:
  - A. Patient has tried and had an inadequate response with a maximally tolerated angiotensin-converting-enzyme inhibitor (ACEI) [e.g., benazepril, lisinopril], or angiotensin II blocker (ARB) [e.g., losartan], or a combination medication containing an ACEI or ARB OR
  - B. Patient has an intolerance or hypersensitivity to an ACEI or ARB, or a combination medication containing an ACEI or ARB, that is not expected to occur with the requested agent OR
  - C. Patient has an FDA labeled contraindication to an ACEI or ARB, or a combination medication containing an ACEI or ARB, that is not expected to occur with the requested agent

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of primary immunoglobulin A nephropathy (IgAN) AND
- 3. The requested agent will be used to reduce proteinuria AND
- 4. Patient has had clinical benefit with the requested agent

### **Age Restriction:**

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., nephrologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Filsuvez PA

## Drug Name(s)

Filsuvez

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. The requested agent will be used for the treatment of wounds associated with dystrophic or junctional epidermolysis bullosa AND
- 2. Patient's diagnosis was confirmed by ONE of the following:
  - A. Immunofluorescence mapping (IFM) OR
  - B. Transmission electron microscopy (TEM) OR
  - C. Genetic testing AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. The requested agent will be used for the treatment of wounds associated with dystrophic or junctional epidermolysis bullosa AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., dermatologist, geneticist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be 3 months for initial, 12 months for renewal

Fintepla PA

## Drug Name(s)

Fintepla

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of seizures associated with Dravet syndrome (DS) or Lennox-Gastaut syndrome (LGS) AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. ALL of the following:
    - i. An echocardiogram assessment will be obtained before and during treatment with the requested agent, to evaluate for valvular heart disease and pulmonary arterial hypertension AND
    - ii. Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis AND
    - iii. Patient does NOT have any FDA labeled contraindications to the requested agent

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Firdapse PA

## Drug Name(s)

Firdapse

### **Indications:**

All FDA-Approved Indications.

#### **Off-Label Uses:**

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require the following:

- 1. Patient has a diagnosis of Lambert-Eaton myasthenic syndrome (LEMS) confirmed by at least ONE of the following:
- A. Electrodiagnostic studies (e.g., electromyography) OR
- B. Antibody testing

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Lambert-Eaton myasthenic syndrome (LEMS) AND
- 3. Patient has had clinical benefit with the requested agent

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Flucytosine PA

### Drug Name(s)

Ancobon

Flucytosine

### **Indications:**

All Medically-Accepted Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. The requested agent will be used in combination with amphotericin B OR
  - B. Prescriber has provided information in support of therapy without concurrent amphotericin B for the requested indication AND
- 3. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

## Age Restriction:

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 10 weeks

Focalin PA

## Drug Name(s)

Dexmethylphenidate Hcl (Focalin)

Focalin

### Indications:

All FDA-Approved Indications.

**Off-Label Uses:** 

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Focalin XR PA

## Drug Name(s)

Dexmethylphenidate Hcl Er (Focalin Xr)

Focalin Xr

### Indications:

All FDA-Approved Indications.

**Off-Label Uses:** 

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Galafold PA

Drug Name(s)

Galafold

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of Fabry disease AND
- 2. The diagnosis was confirmed by mutation of alpha-galactosidase A (alpha-gal A) gene AND
- 3. Patient has an amenable galactosidase alpha gene (GLA) variant mutation AND
- 4. Prescriber has evaluated at least ONE of the following: kidney function (e.g., proteinuria, glomerular filtration rate [GFR]), cardiac function (e.g., left ventricular hypertrophy, conduction defects, mitral and/or aortic valve abnormalities), ophthalmological signs (e.g., corneal verticillate, subcapsular cataracts, conjunctival and/or retinal vasculopathy), peripheral nerve symptoms (e.g., neuropathic pain, heat and/or cold intolerance, impaired sweat function), or gastrointestinal involvement (e.g., nausea, vomiting, abdominal pain, diarrhea, and/or constipation)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Fabry disease AND
- 3. Patient has had clinical benefit with the requested agent [e.g., improvement or stabilization of at least ONE of the following: kidney function (e.g., proteinuria, glomerular filtration rate [GFR]), cardiac function (e.g., left ventricular hypertrophy, conduction defects, mitral and/or aortic valve abnormalities), ophthalmological signs (e.g., corneal verticillate, subcapsular cataracts, conjunctival and/or retinal vasculopathy), peripheral nerve symptoms (e.g., neuropathic pain, heat and/or cold intolerance, impaired sweat function), or gastrointestinal involvement (e.g., nausea, vomiting, abdominal pain, diarrhea, and/or constipation)]

### Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., endocrinologist, geneticist, nephrologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Gamastan PA

### Drug Name(s)

Gamastan

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for approval require ONE of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Primary immunodeficiency [e.g., congenital agammaglobulinemia, common variable immunodeficiency (CVID), severe combined immunodeficiency, Wiskott-Aldrich Syndrome, X-linked agammaglobulinemia (XLA), humoral immunodeficiency, IgG subclass deficiency with or without IgA deficiency] OR
  - B. Multiple sclerosis (MS) AND BOTH of the following:
    - i. Patient has a diagnosis of relapsing remitting MS (RRMS) AND
    - ii. Patient has had an insufficient response, documented failure, or FDA labeled contraindication to TWO MS agents (e.g., Avonex, Betaseron, Copaxone, dimethyl fumarate, fingolimod, glatiramer, Glatopa, Kesimpta, Plegridy, teriflunomide, Vumerity) OR
  - C. Hepatitis A infection prophylaxis AND exposure occurred within the past 2 weeks OR
  - D. Measles (rubeola) prophylaxis AND BOTH of the following:
    - i. Patient is considered susceptible to infection (a susceptible person is defined as one who has not been vaccinated and has not had measles previously) AND
    - ii. Patient was exposed to measles (rubeola) within the past 6 days OR
  - E. Passive immunization against varicella AND BOTH of the following:
    - i. Patient is immunocompromised AND
    - ii. Varicella-Zoster immune globulin is unavailable OR
  - F. Rubella prophylaxis in exposed pregnant woman AND the patient is considered susceptible to infection (a susceptible person is defined as one who has not been vaccinated and has not had rubella previously) OR
- 2. ONE of the following:
  - A. Patient has another FDA labeled indication for the requested agent OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

For prophylaxis diagnoses: see Other Criteria, for all other diagnoses 12 months

## Other Criteria:

Prophylaxis indication with 3 months approval: Hepatitis A infection prophylaxis

Prophylaxis indications with 1 month approval: measles (rubeola) prophylaxis, passive immunization against varicella, rubella prophylaxis in exposed pregnant woman

Gammagard/Gammaked/Gamunex-C PA

# Drug Name(s)

Gammagard Liquid

Gammagard S/D Iga Less Than 1Mcg/Ml

Gammaked

Gamunex-C

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Gattex PA

## Drug Name(s)

Gattex

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of short bowel syndrome (SBS) AND
- 2. Patient is dependent on parenteral nutrition OR intravenous (PN/IV) fluids AND
- 3. ONE of the following:
  - A. Patient is aged 1 year to 17 years AND BOTH of the following:
    - i. A fecal occult blood test has been performed within 6 months prior to initiating treatment with the requested agent AND
    - ii. ONE of the following:
      - a. There was no unexplained blood in the stool OR
      - b. There was unexplained blood in the stool AND a colonoscopy or a sigmoidoscopy was performed OR
  - B. Patient is 18 years of age or over AND BOTH of the following:
    - i. Patient has had a colonoscopy within 6 months prior to initiating treatment with the requested agent AND
    - ii. If polyps were present at this colonoscopy, the polyps were removed AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of short bowel syndrome (SBS) AND
- 3. Patient has had a reduction from baseline in parenteral nutrition OR intravenous (PN/IV) fluids AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be 6 months for initial, 12 months for renewal

Gaucher Enzyme Replacement PA – Cerezyme

#### Drug Name(s)

Cerezyme

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of Gaucher disease type 1 (GD1) confirmed by ONE of the following:
  - A. A baseline (prior to therapy for the requested indication) glucocerebrosidase enzyme activity of less than or equal to 15% of mean normal in peripheral blood leukocytes, fibroblasts, or other nucleated cells OR
  - B. Confirmation of genetic mutation of glucocerebrosidase (GBA) gene with two disease-causing alleles AND
- 2. Prescriber has drawn baseline (prior to therapy for the requested indication) measurements of hemoglobin, platelet count, liver volume, and spleen volume AND
- 3. Prior to any treatment for the intended diagnosis, the patient has had at least ONE of the following clinical presentations:
  - A. Anemia [defined as mean hemoglobin (Hb) level below the testing laboratory's lower limit of the normal range based on age and gender] OR
  - B. Thrombocytopenia (defined as a platelet count of less than 100,000 per microliter) OR
  - C. Hepatomegaly OR
  - D. Splenomegaly OR
  - E. Growth failure (i.e., growth velocity below the standard mean for age) OR
  - F. Evidence of bone disease with other causes ruled out AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Gaucher disease type 1 (GD1) AND
- 3. Patient has had improvement and/or stabilization from baseline (prior to therapy for the requested indication) in at least ONE of the following:
  - A. Hemoglobin (Hb) level OR
  - B. Platelet count OR
  - C. Liver volume OR
  - D. Spleen volume OR
  - E. Growth velocity OR
  - F. Bone pain or disease AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., endocrinologist, geneticist, hematologist, specialist in metabolic diseases) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

**Coverage Duration:** 

Approval will be for 12 months

Gaucher Enzyme Replacement PA - Elelyso

## Drug Name(s)

Elelyso

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of Gaucher disease type 1 (GD1) confirmed by ONE of the following:
  - A. A baseline (prior to therapy for the requested indication) glucocerebrosidase enzyme activity of less than or equal to 15% of mean normal in peripheral blood leukocytes, fibroblasts, or other nucleated cells OR
  - B. Confirmation of genetic mutation of glucocerebrosidase (GBA) gene with two disease-causing alleles AND
- 2. Prescriber has drawn baseline (prior to therapy for the requested indication) measurements of hemoglobin, platelet count, liver volume, and spleen volume AND
- 3. Prior to any treatment for the intended diagnosis, the patient has had at least ONE of the following clinical presentations:
  - A. Anemia [defined as mean hemoglobin (Hb) level below the testing laboratory's lower limit of the normal range based on age and gender] OR
  - B. Thrombocytopenia (defined as a platelet count of less than 100,000 per microliter) OR
  - C. Hepatomegaly OR
  - D. Splenomegaly OR
  - E. Growth failure (i.e., growth velocity below the standard mean for age) OR
  - F. Evidence of bone disease with other causes ruled out AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Gaucher disease type 1 (GD1) AND
- 3. Patient has had improvement and/or stabilization from baseline (prior to therapy for the requested indication) in at least ONE of the following:
  - A. Hemoglobin (Hb) level OR
  - B. Platelet count OR
  - C. Liver volume OR
  - D. Spleen volume OR
  - E. Growth velocity OR
  - F. Bone pain or disease AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., endocrinologist, geneticist, hematologist, specialist in metabolic diseases) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

**Coverage Duration:** 

Approval will be for 12 months

Gaucher Enzyme Replacement PA - Vpriv

# Drug Name(s)

Vpriv

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of Gaucher disease type 1 (GD1) confirmed by ONE of the following:
  - A. A baseline (prior to therapy for the requested indication) glucocerebrosidase enzyme activity of less than or equal to 15% of mean normal in peripheral blood leukocytes, fibroblasts, or other nucleated cells OR
  - B. Confirmation of genetic mutation of glucocerebrosidase (GBA) gene with two disease-causing alleles AND
- 2. Prescriber has drawn baseline (prior to therapy for the requested indication) measurements of hemoglobin, platelet count, liver volume, and spleen volume AND
- 3. Prior to any treatment for the intended diagnosis, the patient has had at least ONE of the following clinical presentations:
  - A. Anemia [defined as mean hemoglobin (Hb) level below the testing laboratory's lower limit of the normal range based on age and gender] OR
  - B. Thrombocytopenia (defined as a platelet count of less than 100,000 per microliter) OR
  - C. Hepatomegaly OR
  - D. Splenomegaly OR
  - E. Growth failure (i.e., growth velocity below the standard mean for age) OR
  - F. Evidence of bone disease with other causes ruled out AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Gaucher disease type 1 (GD1) AND
- 3. Patient has had improvement and/or stabilization from baseline (prior to therapy for the requested indication) in at least ONE of the following:
  - A. Hemoglobin (Hb) level OR
  - B. Platelet count OR
  - C. Liver volume OR
  - D. Spleen volume OR
  - E. Growth velocity OR
  - F. Bone pain or disease AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., endocrinologist, geneticist, hematologist, specialist in metabolic diseases) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

**Coverage Duration:** 

Approval will be for 12 months

Gauze Pads PA

# Drug Name(s)

Gauze Pads

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

This program will be implemented as a dynamic PA.

Criteria for approval require BOTH of the following:

- 1. The requested medical supply product will be used in the delivery of insulin to the body AND
- 2. Patient's medication history includes use of insulin within the past 180 days

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Gralise PA

# Drug Name(s)

Gabapentin Once-Daily

Gralise

### **Indications:**

All FDA-Approved Indications.

## Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of Postherpetic Neuralgia (PHN) AND
- 2. ONE of the following:
  - A. Patient has tried and had an inadequate response to immediate-release gabapentin OR
  - B. Patient has an intolerance or hypersensitivity to immediate-release gabapentin OR
  - C. Patient has an FDA labeled contraindication to immediate-release gabapentin that is not expected to occur with the requested agent

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Postherpetic Neuralgia (PHN) AND
- 3. Patient has had clinical benefit with the requested agent

## Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Growth Hormone PA – Genotropin

# Drug Name(s)

Genotropin

Genotropin Miniquick

### **Indications:**

All Medically-Accepted Indications.

## **Off-Label Uses:**

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

For Children – Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of Turner Syndrome OR
  - B. Patient has a diagnosis of Prader-Willi Syndrome OR
  - C. Patient has a diagnosis of panhypopituitarism AND BOTH of the following:
    - i. Deficiencies in 3 or more pituitary axes AND
    - ii. Measured serum IGF-1 (insulin-like growth factor-1) levels are below the age and sexappropriate reference range when off GH therapy OR
  - D. Patient has a diagnosis of growth hormone deficiency (GHD) or short stature AND BOTH of the following:
    - i. ONE of the following:
      - a. Height more than 2 standard deviations (SD) below the mean for age and sex  $\ensuremath{\mathsf{OR}}$
      - b. Height more than 1.5 SD below the midparental height OR
      - c. A decrease in height SD of more than 0.5 over one year in children at least 2 years of age OR
      - d. Height velocity more than 2 SD below the mean over one year or more than
      - 1.5 SD sustained over two years AND
    - ii. Failure of at least 2 GH stimulation tests (e.g., peak GH value of less than 10 mcg/L after stimulation, or otherwise considered abnormal as determined by testing lab) OR
  - E. Patient has a diagnosis of small for gestational age (SGA) AND ALL of the following:
    - i. Patient is at least 2 years of age AND
    - ii. Documented birth weight and/or length that is 2 or more standard deviations (SD) below the mean for gestational age AND
    - iii. At 24 months of age, the patient fails to manifest catch-up growth evidenced by a height that remains 2 or more SD below the mean for age and sex AND
- 2. ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

## Age Restriction:

## **Prescriber Restrictions:**

**Coverage Duration:** 

# Approval will be for 12 months

# Other Criteria:

For Children – Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Growth Hormone Deficiency, Short Stature OR
  - B. Panhypopituitarism OR
  - C. Prader-Willi Syndrome OR
  - D. Small for Gestational Age OR
  - E. Turner Syndrome AND
- 3. ALL of the following:
  - A. Patient does NOT have closed epiphyses AND
  - B. Patient is being monitored for adverse effects of therapy with the requested agent AND
  - C. Patient's height has increased or height velocity has improved since initiation or last approval of the requested agent

For Adults – Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of childhood growth hormone deficiency (GHD) with genetic or organic origin AND ONE of the following:
    - i. Low IGF-1 (insulin-like growth factor-1) level without GH replacement therapy OR
    - ii. Failure of at least one growth hormone (GH) stimulation test as an adult (e.g., peak GH value of 5 mcg/L or lower after stimulation, or otherwise considered abnormal as determined by testing lab) OR
  - B. Patient has a diagnosis of acquired adult GHD secondary to structural lesions or trauma AND ONE of the following:
    - i. Patient has a diagnosis of panhypopituitarism AND BOTH of the following:
      - a. Deficiencies in 3 or more pituitary axes AND
      - b. Low IGF-1 level without GH replacement therapy OR
    - ii. Patient has failed at least one GH stimulation test as an adult OR
  - C. Patient has a diagnosis of idiopathic GHD (adult or childhood onset) AND the patient has failed at least two growth hormone (GH) stimulation tests as an adult AND
- 2. ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Childhood growth hormone deficiency (GHD) with genetic or organic origin OR
  - B. Acquired adult GHD secondary to structural lesions or trauma OR

- C. Idiopathic GHD (adult or childhood onset) AND
- 3. Patient is being monitored for adverse effects of therapy with the requested agent AND
- 4. Patient's IGF-1 level has been evaluated to confirm the appropriateness of the current dose AND
- 5. Patient has had clinical benefit with the requested agent (i.e., body composition, hip-to-waist ratio, cardiovascular health, bone mineral density, serum cholesterol, physical strength, or quality of life)

Growth Hormone PA – Humatrope

## Drug Name(s)

Humatrope

### **Indications:**

All Medically-Accepted Indications.

## Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

For Children – Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of Turner Syndrome\* OR
  - B. Patient has a diagnosis of SHOX gene deficiency OR
  - C. Patient has a diagnosis of panhypopituitarism\* AND BOTH of the following:
    - i. Deficiencies in 3 or more pituitary axes AND
    - ii. Measured serum IGF-1 (insulin-like growth factor-1) levels are below the age and sexappropriate reference range when off GH therapy OR
  - D. Patient has a diagnosis of growth hormone deficiency (GHD)\* or short stature\* AND BOTH of the following:
    - i. ONE of the following:
      - a. Height more than 2 standard deviation (SD) below the mean for age and sex OR
      - b. Height more than 1.5 SD below the midparental height OR
      - c. A decrease in height SD of more than 0.5 over one year in children at least 2 years of age OR
      - d. Height velocity more than 2 SD below the mean over one year or more than 1.5 SD sustained over 2 years AND
    - ii. Failure of at least 2 GH stimulation tests (e.g., peak GH value of less than 10 mcg/L after stimulation, or otherwise considered abnormal as determined by testing lab) OR
  - E. Patient has a diagnosis of small for gestational age\* AND ALL of the following:
    - i. Patient is at least 2 years of age AND
    - ii. Documented birth weight and/or length that is 2 or more SD below the mean for gestational age AND
    - iii. At 24 months of age, the patient fails to manifest catch-up growth evidenced by a height that remains 2 or more SD below the mean for age and sex AND
- 2. Patient's diagnosis is indicated in the preferred GH agent AND ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

### NOTE:

\*Use of the preferred agent [Omnitrope] is required

NO prerequisites are required for the diagnosis of SHOX gene deficiency

Age Restriction:

**Prescriber Restrictions:** 

## **Coverage Duration:**

Approval will be for 12 months

## Other Criteria:

For Children – Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Growth Hormone Deficiency, Short Stature OR
  - B. Panhypopituitarism OR
  - C. Small for Gestational Age OR
  - D. SHOX gene deficiency OR
  - E. Turner Syndrome AND
- 3. ALL of the following:
  - A. Patient does NOT have closed epiphyses AND
  - B. Patient is being monitored for adverse effects of therapy with the requested agent AND
  - C. Patient's height has increased or height velocity has improved since initiation or last approval of the requested agent

For Adults – Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of childhood growth hormone deficiency (GHD) with genetic or organic origin AND ONE of the following:
    - i. Low IGF-1 (insulin-like growth factor-1) level without GH replacement therapy OR
    - ii. Failure of at least one growth hormone (GH) stimulation test as an adult (e.g., peak GH value of 5 mcg/L or lower after stimulation, or otherwise considered abnormal as determined by testing lab) OR
  - B. Patient has a diagnosis of acquired adult GHD secondary to structural lesions or trauma AND ONE of the following:
    - i. Patient has a diagnosis of panhypopituitarism AND BOTH of the following:
      - a. Deficiencies in 3 or more pituitary axes AND
      - b. Low IGF-1 level without GH replacement therapy OR
    - ii. Patient has failed at least one growth hormone (GH) stimulation test as an adult OR
  - C. Patient has a diagnosis of idiopathic GHD (adult or childhood onset) AND the patient has failed at least two GH stimulation tests as an adult AND
- 2. ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

For Adults – Criteria for renewal approval require ALL of the following:

1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND

- 2. Patient has been diagnosed with ONE of the following:
  - A. Childhood growth hormone deficiency (GHD) with genetic or organic origin OR
  - B. Acquired adult GHD secondary to structural lesions or trauma OR
  - C. Idiopathic GHD (adult or childhood onset) AND
- 3. Patient is being monitored for adverse effects of therapy with the requested agent AND
- 4. Patient's IGF-1 level has been evaluated to confirm the appropriateness of the current dose AND
- 5. Patient has had clinical benefit with the requested agent (i.e., body composition, hip-to-waist ratio, cardiovascular health, bone mineral density, serum cholesterol, physical strength, or quality of life)

Growth Hormone PA – Ngenla

# Drug Name(s)

**Indications:** 

Ngenla

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

For Children – Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of growth hormone deficiency (GHD), short stature, or growth failure due to inadequate secretion of endogenous growth hormone AND BOTH of the following:
  - A. ONE of the following:
    - i. Height more than 2 standard deviations (SD) below the mean for age and sex OR
    - ii. Height more than 1.5 SD below the midparental height OR
    - iii. A decrease in height SD of more than 0.5 over one year in children at least 2 years of age OR
    - iv. Height velocity more than 2 SD below the mean over one year or more than 1.5 SD sustained over two years AND
  - B. Failure of at least 2 GH stimulation tests (e.g., peak GH value of less than 10 mcg/L after stimulation, or otherwise considered abnormal as determined by testing lab) AND
- 2. ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

For Children – Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of growth hormone deficiency (GHD), short stature, or growth failure due to inadequate secretion of endogenous growth hormone AND
- 3. ALL of the following:
  - A. Patient does NOT have closed epiphyses AND
  - B. Patient is being monitored for adverse effects of therapy with the requested agent AND
  - C. Patient's height has increased or height velocity has improved since initiation or last approval of the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Growth Hormone PA – Norditropin

# Drug Name(s)

Norditropin Flexpro

### **Indications:**

All Medically-Accepted Indications.

## Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

For Children – Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of Turner Syndrome\* OR
  - B. Patient has a diagnosis of Prader-Willi Syndrome\* OR
  - C. Patient has a diagnosis of Noonan Syndrome OR
  - D. Patient has a diagnosis of panhypopituitarism\* AND BOTH of the following:
    - i. Deficiencies in 3 or more pituitary axes AND
    - ii. Measured serum IGF-1 (insulin-like growth factor-1) levels are below the age and sexappropriate reference range when off GH therapy OR
  - E. Patient has a diagnosis of growth hormone deficiency (GHD)\* or short stature\* AND BOTH of the following:
    - i. ONE of the following:
      - a. Height more than 2 standard deviations (SD) below the mean for age and sex OR
      - b. Height more than 1.5 SD below the midparental height OR
      - c. A decrease in height SD of more than 0.5 over one year in children at least 2 years of age OR
      - d. Height velocity more than 2 SD below the mean over one year or more than 1.5 SD sustained over 2 years AND
    - ii. Failure of at least 2 GH stimulation tests (e.g., peak GH value of less than 10 mcg/L after stimulation, or otherwise considered abnormal as determined by testing lab) OR
  - F. Patient has a diagnosis of small for gestational age\* AND ALL of the following:
    - i. Patient is at least 2 years of age AND
    - ii. Documented birth weight and/or length that is 2 or more SD below the mean for gestational age AND
    - iii. At 24 months of age, the patient fails to manifest catch-up growth evidenced by a height that remains 2 or more SD below the mean for age and sex AND
- 2. Patient's diagnosis is indicated in the preferred GH agent AND ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

## NOTE:

\*Use of the preferred agent [Omnitrope] is required

NO prerequisites are required for the diagnosis of Noonan Syndrome

# Age Restriction:

## **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

For Children – Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Growth Hormone Deficiency (GHD), Short Stature OR
  - B. Noonan Syndrome OR
  - C. Panhypopituitarism OR
  - D. Small for Gestational Age OR
  - E. Turner Syndrome OR
  - F. Prader-Willi Syndrome AND
- 3. ALL of the following:
  - A. Patient does NOT have closed epiphyses AND
  - B. Patient is being monitored for adverse effects of therapy with the requested agent AND
  - C. Patient's height has increased or height velocity has improved since initiation or last approval of the requested agent

For Adults – Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of childhood GHD with genetic or organic origin AND ONE of the following:
    - i. Low IGF-1 (insulin-like growth factor-1) level without GH replacement therapy OR
    - ii. Failure of at least one growth hormone (GH) stimulation test as an adult (e.g., peak GH value of 5 mcg/L or lower after stimulation, or otherwise considered abnormal as determined by testing lab) OR
  - B. Patient has a diagnosis of acquired adult GHD secondary to structural lesions or trauma AND ONE of the following:
    - i. Patient has a diagnosis of panhypopituitarism AND BOTH of the following:
    - a. Deficiencies in 3 or more pituitary axes AND
    - b. Low IGF-1 level without GH replacement therapy OR
    - ii. Patient has failed at least one growth hormone (GH) stimulation test as an adult OR
  - C. Patient has a diagnosis of idiopathic GHD (adult or childhood onset) AND the patient has failed at least two GH stimulation tests as an adult AND
- 2. ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Childhood growth hormone deficiency (GHD) with genetic or organic origin OR
  - B. Acquired adult GHD secondary to structural lesions or trauma OR
  - C. Idiopathic GHD (adult or childhood onset) AND
- 3. Patient is being monitored for adverse effects of therapy with the requested agent AND
- 4. Patient's IGF-1 level has been evaluated to confirm the appropriateness of the current dose AND
- 5. Patient has had clinical benefit with the requested agent (i.e., body composition, hip-to-waist ratio, cardiovascular health, bone mineral density, serum cholesterol, physical strength, or quality of life)

Growth Hormone PA - Nutropin

# Drug Name(s)

Nutropin Aq Nuspin 10

Nutropin Aq Nuspin 20

Nutropin Aq Nuspin 5

## **Indications:**

All Medically-Accepted Indications.

## Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

For Children – Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of Turner Syndrome\* OR
  - B. Patient has a diagnosis of chronic renal insufficiency AND BOTH of the following:
    - i. Height velocity (HV) is greater than or equal to 1.88 standard deviations (SD) below the mean or less than the third percentile for age and sex AND
    - ii. Other etiologies for growth impairment have been addressed OR
  - C. Patient has a diagnosis of panhypopituitarism\* AND BOTH of the following:
    - i. Deficiencies in 3 or more pituitary axes AND
    - ii. Measured serum insulin-like growth factor-1 levels are below the age and sexappropriate reference range when off GH therapy OR
  - D. Patient has a diagnosis of growth hormone deficiency (GHD)\* or short stature\* AND BOTH of the following:
    - i. ONE of the following:
      - a. Height more than 2 SD below the mean for age and sex OR
      - b. Height more than 1.5 SD below the midparental height OR
      - c. A decrease in height SD of more than 0.5 over one year in children at least 2 years of age OR
      - d. Height velocity more than 2 SD below the mean over one year or more than
      - 1.5 SD sustained over 2 years AND
    - ii. Failure of at least 2 GH stimulation tests (e.g., peak GH value of less than 10 mcg/L after stimulation, or otherwise considered abnormal as determined by testing lab) AND
- 2. Patient's diagnosis is indicated in the preferred GH agent AND ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

### NOTE:

\*Use of the preferred agent [Omnitrope] is required

NO prerequisites are required for the diagnosis of chronic renal insufficiency

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

### Other Criteria:

For Children – Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Growth Hormone Deficiency, Short Stature OR
  - B. Chronic renal insufficiency OR
  - C. Panhypopituitarism OR
  - D. Turner Syndrome AND
- 3. ALL of the following:
  - A. Patient does NOT have closed epiphyses AND
  - B. Patient is being monitored for adverse effects of therapy with the requested agent AND
  - C. Patient's height has increased or height velocity has improved since initiation or last approval of the requested agent

For Adults – Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of childhood growth hormone deficiency (GHD) with genetic or organic origin AND ONE of the following:
    - i. Low IGF-1 (insulin-like growth factor-1) level without GH replacement therapy OR
    - ii. Failure of at least one growth hormone (GH) stimulation test as an adult (e.g., peak GH value of 5 mcg/L or lower after stimulation, or otherwise considered abnormal as determined by testing lab) OR
  - B. Patient has a diagnosis of acquired adult GHD secondary to structural lesions or trauma AND ONE of the following:
    - i. Patient has a diagnosis of panhypopituitarism AND BOTH of the following:
      - a. Deficiencies in 3 or more pituitary axes AND
      - b. Low IGF-1 level without GH replacement therapy OR
    - ii. Patient has failed at least one growth hormone (GH) stimulation test as an adult OR
  - C. Patient has a diagnosis of idiopathic GHD (adult or childhood onset) AND the patient has failed at least two growth hormone (GH) stimulation tests as an adult AND
- 2. ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Childhood growth hormone deficiency (GHD) with genetic or organic origin OR
  - B. Acquired adult GHD secondary to structural lesions or trauma OR

- C. Idiopathic GHD (adult or childhood onset) AND
- 3. Patient is being monitored for adverse effects of therapy with the requested agent AND
- 4. Patient's IGF-1 level has been evaluated to confirm the appropriateness of the current dose AND
- 5. Patient has had clinical benefit with the requested agent (i.e., body composition, hip-to-waist ratio, cardiovascular health, bone mineral density, serum cholesterol, physical strength, or quality of life)

Growth Hormone PA – Omnitrope

# Drug Name(s)

Omnitrope

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

For Children – Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of Turner Syndrome OR
  - B. Patient has a diagnosis of Prader-Willi Syndrome OR
  - C. Patient has a diagnosis of panhypopituitarism AND BOTH of the following:
    - i. Deficiencies in 3 or more pituitary axes AND
    - ii. Measured serum IGF-1 (insulin-like growth factor-1) levels are below the age and sexappropriate reference range when off GH therapy OR
  - D. Patient has a diagnosis of growth hormone deficiency (GHD) or short stature AND BOTH of the following:
    - i. Patient has ONE of the following:
      - a. Height more than 2 standard deviations (SD) below the mean for age and sex OR
      - b. Height more than 1.5 SD below the midparental height OR
      - c. A decrease in height SD of more than 0.5 over one year in children at least 2 years of age OR
      - d. Height velocity more than 2 SD below the mean over one year or more than 1.5 SD sustained over two years AND
    - ii. Failure of at least 2 growth hormone (GH) stimulation tests (e.g., peak GH value of less than 10 mcg/L after stimulation, or otherwise considered abnormal as determined by testing lab) OR
  - E. Patient has a diagnosis of small for gestational age (SGA) AND ALL of the following:
    - i. Patient is at least 2 years of age AND
    - ii. Documented birth weight and/or length that is 2 or more SD below the mean for gestational age AND
    - iii. At 24 months of age, the patient fails to manifest catch-up growth evidenced by a height that remains 2 or more SD below the mean for age and sex

# Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

- 1. Patient has been previously approved for the preferred agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Growth Hormone Deficiency, Short Stature OR
  - B. Panhypopituitarism OR
  - C. Prader-Willi Syndrome OR
  - D. Small for Gestational Age (SGA) OR
  - E. Turner Syndrome AND
- 3. ALL of the following:
  - A. Patient does NOT have closed epiphyses AND
  - B. Patient is being monitored for adverse effects of therapy with the requested agent AND
  - C. Patient's height has increased or height velocity has improved since initiation or last approval of the requested agent

For Adults – Criteria for initial approval require the following:

- 1. Patient has been diagnosed with ONE of the following:
  - A. Childhood growth hormone deficiency (GHD) with genetic or organic origin AND ONE of the following:
    - i. Low IGF-1 (insulin-like growth factor-1) level without GH replacement therapy OR
    - ii. Failure of at least one growth hormone (GH) stimulation test as an adult (e.g., peak GH value of 5 mcg/L or lower after stimulation, or otherwise considered abnormal as determined by testing lab) OR
  - B. Acquired adult GHD secondary to structural lesions or trauma AND ONE of the following:
    - i. Patient has a diagnosis of panhypopituitarism AND BOTH of the following:
      - a. Deficiencies in 3 or more pituitary axes AND
      - b. Low IGF-1 level without GH replacement therapy OR
  - ii. Patient has failed at least one growth hormone (GH) stimulation test as an adult OR C. Idiopathic GHD (adult or childhood onset) AND the patient has failed at least two growth hormone (GH) stimulation tests as an adult

- 1. Patient has been previously approved for the preferred agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Childhood growth hormone deficiency (GHD) with genetic or organic origin OR
  - B. Acquired adult GHD secondary to structural lesions or trauma OR
  - C. Idiopathic GHD (adult or childhood onset) AND
- 3. Patient is being monitored for adverse effects of therapy with the requested agent AND
- 4. Patient's IGF-1 level has been evaluated to confirm the appropriateness of the current dose AND
- 5. Patient has had clinical benefit with the requested agent (i.e., body composition, hip-to-waist ratio, cardiovascular health, bone mineral density, serum cholesterol, physical strength, or quality of life)

Growth Hormone PA – Serostim

# Drug Name(s)

Serostim

### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. The requested agent is for the treatment of an HIV patient with wasting or cachexia AND
- 2. BOTH of the following:
  - A. The requested agent will be used in combination with antiretroviral therapy AND
  - B. ONE of the following:
    - i. Patient has had an unintentional weight loss of 10% or more of body weight over 12 months OR
    - ii. Patient has had an unintentional weight loss of greater than 7.5% over 6 months OR
    - iii. Patient has a mid-upper arm circumference less than 10th percentile OR
    - iv. Patient has a body cell mass (BCM) loss of 5% or more over 6 months OR
    - v. Patient's sex is male, has a BCM less than 35% AND a BMI of less than 27 kg/m2 OR
    - vi. Patient's sex is female, has a BCM less than 23% AND a BMI of less than 27 kg/m2

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. The requested agent is for the treatment of an HIV patient with wasting or cachexia AND
- 3. BOTH of the following:
  - A. The requested agent will be used in combination with antiretroviral therapy AND
  - B. Patient has had clinical benefit with the requested agent (e.g., weight increase or weight stabilization)

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 weeks

Growth Hormone PA – Skytrofa

# Drug Name(s)

Skytrofa

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

For Children – Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of growth hormone deficiency (GHD), short stature, or growth failure due to inadequate secretion of endogenous growth hormone AND BOTH of the following:
  - A. ONE of the following:
    - i. Height more than 2 standard deviations (SD) below the mean for age and sex OR
    - ii. Height more than 1.5 SD below the midparental height OR
    - iii. A decrease in height SD of more than 0.5 over one year in children at least 2 years of age OR
    - iv. Height velocity more than 2 SD below the mean over one year or more than 1.5 SD sustained over two years AND
  - B. Failure of at least 2 GH stimulation tests (e.g., peak GH value of less than 10 mcg/L after stimulation, or otherwise considered abnormal as determined by testing lab) AND
- 2. ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

For Children – Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of growth hormone deficiency (GHD), short stature, or growth failure due to inadequate secretion of endogenous growth hormone AND
- 3. ALL of the following:
  - A. Patient does NOT have closed epiphyses AND
  - B. Patient is being monitored for adverse effects of therapy with the requested agent AND
  - C. Patient's height has increased or height velocity has improved since initiation or last approval of the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Growth Hormone PA - Sogroya

## Drug Name(s)

Sogroya

## **Indications:**

All Medically-Accepted Indications.

### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

For Children – Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of growth hormone deficiency (GHD), short stature, or growth failure due to inadequate secretion of endogenous growth hormone AND BOTH of the following:
  - A. ONE of the following:
    - i. Height more than 2 standard deviations (SD) below the mean for age and sex OR
    - ii. Height more than 1.5 SD below the midparental height OR
    - iii. A decrease in height SD of more than 0.5 over one year in children at least 2 years of age OR
    - iv. Height velocity more than 2 SD below the mean over one year or more than 1.5 SD sustained over two years AND
  - B. Failure of at least 2 GH stimulation tests (e.g., peak GH value of less than 10 mcg/L after stimulation, or otherwise considered abnormal as determined by testing lab) AND
- 2. ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

For Children – Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of growth hormone deficiency (GHD), short stature, or growth failure due to inadequate secretion of endogenous growth hormone AND
- 3. ALL of the following:
  - A. Patient does NOT have closed epiphyses AND
  - B. Patient is being monitored for adverse effects of therapy with the requested agent AND
  - C. Patient's height has increased or height velocity has improved since initiation or last approval of the requested agent

# Age Restriction:

## **Prescriber Restrictions:**

## **Coverage Duration:**

Approval will be for 12 months

## Other Criteria:

For Adults – Criteria for initial approval require BOTH of the following:

1. ONE of the following:

- A. Patient has a diagnosis of childhood growth hormone deficiency (GHD) with genetic or organic origin AND ONE of the following:
  - i. Low IGF-1 (insulin-like growth factor-1) level without GH replacement therapy OR ii. Failure of at least one growth hormone (GH) stimulation test as an adult (e.g., peak GH value of 5 mcg/L or lower after stimulation, or otherwise considered abnormal as determined by testing lab) OR
- B. Patient has a diagnosis of acquired adult GHD secondary to structural lesions or trauma AND ONE of the following:
  - i. Patient has a diagnosis of panhypopituitarism AND BOTH of the following:
    - a. Deficiencies in 3 or more pituitary axes AND
    - b. Low IGF-1 level without GH replacement therapy OR
  - ii. Patient has failed at least one growth hormone (GH) stimulation test as an adult OR
- C. Patient has a diagnosis of idiopathic GHD (adult or childhood onset) AND the patient has failed at least two GH stimulation tests as an adult AND
- 2. ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Childhood growth hormone deficiency (GHD) with genetic or organic origin OR
  - B. Acquired adult GHD secondary to structural lesions or trauma OR
  - C. Idiopathic GHD (adult or childhood onset) AND
- 3. Patient is being monitored for adverse effects of therapy with the requested agent AND
- 4. Patient's IGF-1 level has been evaluated to confirm the appropriateness of the current dose AND
- 5. Patient has had clinical benefit with the requested agent (i.e., body composition, hip-to-waist ratio, cardiovascular health, bone mineral density, serum cholesterol, physical strength, or quality of life)

Growth Hormone PA - Zomacton

# Drug Name(s)

Zomacton

## **Indications:**

All Medically-Accepted Indications.

### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

For Children – Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
- A. Patient has a diagnosis of Turner Syndrome\* OR
- B. Patient has a diagnosis of SHOX gene deficiency OR
- C. Patient has a diagnosis of panhypopituitarism\* AND BOTH of the following:
- i. Deficiencies in 3 or more pituitary axes AND
- ii. Measured serum IGF-1 (insulin-like growth factor-1) levels are below the age and sex-appropriate reference range when off GH therapy OR
- D. Patient has a diagnosis of growth hormone deficiency (GHD)\* or short stature\* AND BOTH of the following:
- i. ONE of the following:
- a. Height more than 2 standard deviation (SD) below the mean for age and sex OR
- b. Height more than 1.5 SD below the midparental height OR
- c. A decrease in height SD of more than 0.5 over one year in children at least 2 years of age OR
- d. Height velocity more than 2 SD below the mean over one year or more than 1.5 SD sustained over 2 years AND
- ii. Failure of at least 2 GH stimulation tests (e.g., peak GH value of less than 10 mcg/L after stimulation, or otherwise considered abnormal as determined by testing lab) OR
- E. Patient has a diagnosis of small for gestational age\* AND ALL of the following:
- i. Patient is at least 2 years of age AND
- ii. Documented birth weight and/or length that is 2 or more SD below the mean for gestational age AND
- iii. At 24 months of age, the patient fails to manifest catch-up growth evidenced by a height that remains 2 or more SD below the mean for age and sex AND
- 2. Patient's diagnosis is indicated in the preferred GH agent AND ONE of the following:
- A. Patient has tried and failed the preferred agent [Omnitrope] OR
- B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

### NOTE:

\*Use of the preferred agent [Omnitrope] is required

NO prerequisites are required for the diagnosis of SHOX gene deficiency

# Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

# Approval will be for 12 months

# Other Criteria:

For Children – Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Growth Hormone Deficiency, Short Stature OR
  - B. Panhypopituitarism OR
  - C. Small for Gestational Age OR
  - D. SHOX gene deficiency OR
  - E. Turner Syndrome AND
- 3. ALL of the following:
  - A. Patient does NOT have closed epiphyses AND
  - B. Patient is being monitored for adverse effects of therapy with the requested agent AND
  - C. Patient's height has increased or height velocity has improved since initiation or last approval of the requested agent

For Adults – Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of childhood growth hormone deficiency (GHD) with genetic or organic origin AND ONE of the following:
    - i. Low IGF-1 (insulin-like growth factor-1) level without GH replacement therapy OR
    - ii. Failure of at least one growth hormone (GH) stimulation test as an adult (e.g., peak GH value of 5 mcg/L or lower after stimulation, or otherwise considered abnormal as determined by testing lab) OR
  - B. Patient has a diagnosis of acquired adult GHD secondary to structural lesions or trauma AND ONE of the following:
    - i. Patient has a diagnosis of panhypopituitarism AND BOTH of the following:
      - a. Deficiencies in 3 or more pituitary axes AND
      - b. Low IGF-1 level without GH replacement therapy OR
    - ii. Patient has failed at least one growth hormone (GH) stimulation test as an adult OR
  - C. Patient has a diagnosis of idiopathic GHD (adult or childhood onset) AND the patient has failed at least two GH stimulation tests as an adult AND
- 2. ONE of the following:
  - A. Patient has tried and failed the preferred agent [Omnitrope] OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to the preferred agent [Omnitrope]

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has been diagnosed with ONE of the following:
  - A. Childhood growth hormone deficiency (GHD) with genetic or organic origin OR
  - B. Acquired adult GHD secondary to structural lesions or trauma OR

- C. Idiopathic GHD (adult or childhood onset) AND
- 3. Patient is being monitored for adverse effects of therapy with the requested agent AND
- 4. Patient's IGF-1 level has been evaluated to confirm the appropriateness of the current dose AND
- 5. Patient has had clinical benefit with the requested agent (i.e., body composition, hip-to-waist ratio, cardiovascular health, bone mineral density, serum cholesterol, physical strength, or quality of life)

HAE PA - Berinert

## Drug Name(s)

Berinert

### Indications:

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of hereditary angioedema (HAE) which has been confirmed via two confirmatory tests of C1-INH antigenic level, C1-INH functional level, and C4 level as follows:
  - a. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type I)]: decreased quantities of C4 and C1-INH (antigenic and functional level) OR
  - b. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type II)]: decreased quantities of C4 and C1-INH function (C1-INH protein level may be normal) OR
  - c. Hereditary angioedema (HAE) with normal C1INH [HAE-nI-C1INH (Type III)]: Normal levels of C4 and C1-INH [antigenic and functional level (at baseline and during an attack)] AND ONE of the following:
    - i. BOTH of the following:
      - 1. Family history of angioedema AND
      - 2. ALL other causes of angioedema have been ruled out OR
    - ii. Patient demonstrates a Factor XII mutation, angiopoietin-1 (ANGPT1) mutation, plasminogen (PLG) mutation, kininogen1 mutation, heparan sulfate 3-O-sulfotransferase 6 gene mutation, or myoferlin gene mutation that is associated with the disease AND
- 2. Medications known to cause angioedema (e.g., ACE-Inhibitors, estrogens, angiotensin II receptor blockers) have been evaluated and discontinued when appropriate AND
- 3. The requested agent will be used to treat acute HAE attacks AND
- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for treatment of acute HAE attacks

## Age Restriction:

## **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

### Other Criteria:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of hereditary angioedema (HAE) AND
- 3. The requested agent will be used to treat acute HAE attacks AND
- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for treatment of acute HAE attacks AND

5. Patient has had a decrease in the frequency or severity of acute attacks or stabilization of disease from use of the requested agent

HAE PA - Cinryze

# Drug Name(s)

Cinryze

### **Indications:**

All FDA-Approved Indications, Some Medically-Accepted Indications.

## **Off-Label Uses:**

Acute HAE attacks

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of hereditary angioedema (HAE) which has been confirmed via two confirmatory tests of C1-INH antigenic level, C1-INH functional level, and C4 level as follows:
  - a. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type I)]: decreased quantities of C4 and C1-INH (antigenic and functional level) OR
  - b. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type II)]: decreased quantities of C4 and C1-INH function (C1-INH protein level may be normal) OR
  - c. Hereditary angioedema (HAE) with normal C1INH [HAE-nI-C1INH (Type III)]: Normal levels of C4 and C1-INH [antigenic and functional level (at baseline and during an attack)] AND ONE of the following:
    - i. BOTH of the following:
      - 1. Family history of angioedema AND
      - 2. ALL other causes of angioedema have been ruled out OR
    - ii. Patient demonstrates a Factor XII mutation, angiopoietin-1 (ANGPT1) mutation, plasminogen (PLG) mutation, kininogen1 mutation, heparan sulfate 3-O-sulfotransferase 6 gene mutation, or myoferlin gene mutation that is associated with the disease AND
- 2. Medications known to cause angioedema (e.g., ACE-Inhibitors, estrogens, angiotensin II receptor blockers) have been evaluated and discontinued when appropriate AND
- 3. ONE of the following:
  - a. The requested agent will be used to treat acute HAE attacks AND the patient will NOT be using the requested agent in combination with another HAE agent indicated for treatment of acute HAE attacks OR
  - b. The requested agent will be used for prophylaxis against HAE attacks AND the patient will NOT be using the requested agent in combination with another HAE agent indicated for prophylaxis against HAE attacks

# Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

## Other Criteria:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of hereditary angioedema (HAE) AND ONE of the following:
  - a. The requested agent will be used to treat acute HAE attacks AND the patient will NOT be using the requested agent in combination with another HAE agent indicated for treatment of acute HAE attacks OR
  - b. The requested agent will be used for prophylaxis against HAE attacks AND the patient will NOT be using the requested agent in combination with another HAE agent indicated for prophylaxis against HAE attacks AND
- 3. Patient has had a decrease in the frequency or severity of acute attacks or has had stabilization of disease from use of the requested agent

HAE PA - Haegarda

## Drug Name(s)

Haegarda

### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of hereditary angioedema (HAE) which has been confirmed via two confirmatory tests of C1-INH antigenic level, C1-INH functional level, and C4 level as follows:
  - a. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type I)]: decreased quantities of C4 and C1-INH (antigenic and functional level) OR
  - b. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type II)]: decreased quantities of C4 and C1-INH function (C1-INH protein level may be normal) OR
  - c. Hereditary angioedema (HAE) with normal C1INH [HAE-nI-C1INH (Type III)]: Normal levels of C4 and C1-INH [antigenic and functional level (at baseline and during an attack)] AND ONE of the following:
    - i. BOTH of the following:
      - 1. Family history of angioedema AND
      - 2. ALL other causes of angioedema have been ruled out OR
    - ii. Patient demonstrates a Factor XII mutation, angiopoietin-1 (ANGPT1) mutation, plasminogen (PLG) mutation, kininogen1 mutation, heparan sulfate 3-O-sulfotransferase 6 gene mutation, or myoferlin gene mutation that is associated with the disease AND
- 2. Medications known to cause angioedema (e.g., ACE-Inhibitors, estrogens, angiotensin II receptor blockers) have been evaluated and discontinued when appropriate AND
- 3. The requested agent will be used for prophylaxis against HAE attacks AND
- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for prophylaxis against HAE attacks

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

## **Prescriber Restrictions:**

### **Coverage Duration:**

Approval will be for 12 months

### Other Criteria:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of hereditary angioedema (HAE) AND
- 3. The requested agent is being used for prophylaxis against HAE attacks AND
- 4. Patient has had a decrease in the frequency or severity of acute attacks or has had stabilization of disease from use of the requested agent AND

5. Patient will NOT be using the requested agent in combination with another HAE agent indicated for prophylaxis against HAE attacks

HAE PA – Icatibant

# Drug Name(s)

Firazyr

Icatibant Acetate

Sajazir

## **Indications:**

All FDA-Approved Indications.

## Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of hereditary angioedema (HAE) which has been confirmed via two confirmatory tests of C1-INH antigenic level, C1-INH functional level, and C4 level as follows:
  - a. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type I)]: decreased quantities of C4 and C1-INH (antigenic and functional level) OR
  - b. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type II)]: decreased quantities of C4 and C1-INH function (C1-INH protein level may be normal) OR
  - c. Hereditary angioedema (HAE) with normal C1INH [HAE-nI-C1INH (Type III)]: Normal levels of C4 and C1-INH [antigenic and functional level (at baseline and during an attack)] AND ONE of the following:
    - i. BOTH of the following:
      - 1. Family history of angioedema AND
      - 2. ALL other causes of angioedema have been ruled out OR
    - ii. Patient demonstrates a Factor XII mutation, angiopoietin-1 (ANGPT1) mutation, plasminogen (PLG) mutation, kininogen1 mutation, heparan sulfate 3-O-sulfotransferase 6 gene mutation, or myoferlin gene mutation that is associated with the disease AND
- 2. Medications known to cause angioedema (e.g., ACE-Inhibitors, estrogens, angiotensin II receptor blockers) have been evaluated and discontinued when appropriate AND
- 3. The requested agent will be used to treat acute HAE attacks AND
- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for treatment of acute HAE attacks

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

# **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

# Other Criteria:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of hereditary angioedema (HAE) AND
- 3. The requested agent will be used to treat acute HAE attacks AND

- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for treatment of acute HAE attacks AND
- 5. Patient has had a decrease in the frequency or severity of acute attacks or stabilization of disease from use of the requested agent

HAE PA – Kalbitor

## Drug Name(s)

Kalbitor

#### Indications:

All FDA-Approved Indications.

### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of hereditary angioedema (HAE) which has been confirmed via two confirmatory tests of C1-INH antigenic level, C1-INH functional level, and C4 level as follows:
  - a. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type I)]: decreased quantities of C4 and C1-INH (antigenic and functional level) OR
  - b. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type II)]: decreased quantities of C4 and C1-INH function (C1-INH protein level may be normal) OR
  - c. Hereditary angioedema (HAE) with normal C1INH [HAE-nI-C1INH (Type III)]: Normal levels of C4 and C1-INH [antigenic and functional level (at baseline and during an attack)] AND ONE of the following:
    - i. BOTH of the following:
      - 1. Family history of angioedema AND
      - 2. ALL other causes of angioedema have been ruled out OR
    - ii. Patient demonstrates a Factor XII mutation, angiopoietin-1 (ANGPT1) mutation, plasminogen (PLG) mutation, kininogen1 mutation, heparan sulfate 3-O-sulfotransferase 6 gene mutation, or myoferlin gene mutation that is associated with the disease AND
- 2. Medications known to cause angioedema (e.g., ACE-Inhibitors, estrogens, angiotensin II receptor blockers) have been evaluated and discontinued when appropriate AND
- 3. The requested agent will be used to treat acute HAE attacks AND
- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for treatment of acute HAE attacks

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

## **Prescriber Restrictions:**

### **Coverage Duration:**

Approval will be for 12 months

### Other Criteria:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of hereditary angioedema (HAE) AND
- 3. The requested agent will be used to treat acute HAE attacks AND
- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for treatment of acute HAE attacks AND

5. Patient has had a decrease in the frequency or severity of acute attacks or stabilization of disease from use of the requested agent	

HAE PA - Orladeyo

## Drug Name(s)

Orladeyo

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of hereditary angioedema (HAE) which has been confirmed via two confirmatory tests of C1-INH antigenic level, C1-INH functional level, and C4 level as follows:
  - a. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type I)]: decreased quantities of C4 and C1-INH (antigenic and functional level) OR
  - b. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type II)]: decreased quantities of C4 and C1-INH function (C1-INH protein level may be normal) OR
  - c. Hereditary angioedema (HAE) with normal C1INH [HAE-nI-C1INH (Type III)]: Normal levels of C4 and C1-INH [antigenic and functional level (at baseline and during an attack)] AND ONE of the following:
    - i. BOTH of the following:
      - 1. Family history of angioedema AND
      - 2. ALL other causes of angioedema have been ruled out OR
    - ii. Patient demonstrates a Factor XII mutation, angiopoietin-1 (ANGPT1) mutation, plasminogen (PLG) mutation, kininogen1 mutation, heparan sulfate 3-O-sulfotransferase 6 gene mutation, or myoferlin gene mutation that is associated with the disease AND
- 2. Medications known to cause angioedema (e.g., ACE-Inhibitors, estrogens, angiotensin II receptor blockers) have been evaluated and discontinued when appropriate AND
- 3. The requested agent will be used for prophylaxis against HAE attacks AND
- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for prophylaxis against HAE attacks

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of hereditary angioedema (HAE) AND
- 3. The requested agent is being used for prophylaxis against HAE attacks AND
- 4. Patient has had a decrease in the frequency or severity of acute attacks or has had stabilization of disease from use of the requested agent AND

5. Patient will NOT be using the requested agent in combination with another HAE agent indicated for prophylaxis against HAE attacks

HAE PA – Ruconest

#### Drug Name(s)

Ruconest

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of hereditary angioedema (HAE) which has been confirmed via two confirmatory tests of C1-INH antigenic level, C1-INH functional level, and C4 level as follows:
  - a. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type I)]: decreased quantities of C4 and C1-INH (antigenic and functional level) OR
  - b. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type II)]: decreased quantities of C4 and C1-INH function (C1-INH protein level may be normal) OR
  - c. Hereditary angioedema (HAE) with normal C1INH [HAE-nI-C1INH (Type III)]: Normal levels of C4 and C1-INH [antigenic and functional level (at baseline and during an attack)] AND ONE of the following:
    - i. BOTH of the following:
      - 1. Family history of angioedema AND
      - 2. ALL other causes of angioedema have been ruled out OR
    - ii. Patient demonstrates a Factor XII mutation, angiopoietin-1 (ANGPT1) mutation, plasminogen (PLG) mutation, kininogen1 mutation, heparan sulfate 3-O-sulfotransferase 6 gene mutation, or myoferlin gene mutation that is associated with the disease AND
- 2. Medications known to cause angioedema (e.g., ACE-Inhibitors, estrogens, angiotensin II receptor blockers) have been evaluated and discontinued when appropriate AND
- 3. The requested agent will be used to treat acute HAE attacks AND
- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for treatment of acute HAE attacks

### Age Restriction:

#### **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of hereditary angioedema (HAE) AND
- 3. The requested agent will be used to treat acute HAE attacks AND
- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for treatment of acute HAE attacks AND

5. Patient has had a decrease in the frequency or severity of acute attacks or stabilization of disease from use of the requested agent

HAE PA - Takhzyro

## Drug Name(s)

Takhzyro

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of hereditary angioedema (HAE) which has been confirmed via two confirmatory tests of C1-INH antigenic level, C1-INH functional level, and C4 level as follows:
  - a. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type I)]: decreased quantities of C4 and C1-INH (antigenic and functional level) OR
  - b. Hereditary angioedema (HAE) due to C1INH deficiency [HAE-C1INH (Type II)]: decreased quantities of C4 and C1-INH function (C1-INH protein level may be normal) OR
  - c. Hereditary angioedema (HAE) with normal C1INH [HAE-nI-C1INH (Type III)]: Normal levels of C4 and C1-INH [antigenic and functional level (at baseline and during an attack)] AND ONE of the following:
    - i. BOTH of the following:
      - 1. Family history of angioedema AND
      - 2. ALL other causes of angioedema have been ruled out OR
    - ii. Patient demonstrates a Factor XII mutation, angiopoietin-1 (ANGPT1) mutation, plasminogen (PLG) mutation, kininogen1 mutation, heparan sulfate 3-O-sulfotransferase 6 gene mutation, or myoferlin gene mutation that is associated with the disease AND
- 2. Medications known to cause angioedema (e.g., ACE-Inhibitors, estrogens, angiotensin II receptor blockers) have been evaluated and discontinued when appropriate AND
- 3. The requested agent will be used for prophylaxis against HAE attacks AND
- 4. Patient will NOT be using the requested agent in combination with another HAE agent indicated for prophylaxis against HAE attacks

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of hereditary angioedema (HAE) AND
- 3. The requested agent is being used for prophylaxis against HAE attacks AND
- 4. Patient has had a decrease in the frequency or severity of acute attacks or has had stabilization of disease from use of the requested agent AND

5. Patient will NOT be using the requested agent in combination with another HAE agent indicated for prophylaxis against HAE attacks

Harvoni PA

## Drug Name(s)

Harvoni

Ledipasvir/Sofosbuvir

#### **Indications:**

All Medically-Accepted Indications.

#### **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of hepatitis C confirmed by serological markers AND
- 2. Prescriber has screened the patient for current or prior hepatitis B viral (HBV) infection and if positive, will monitor the patient for HBV flare-up or reactivation during and after treatment with the requested agent AND
- 3. The requested agent will be used in a treatment regimen and length of therapy that is supported in FDA approved labeling or AASLD/IDSA guidelines for the patient's diagnosis and genotype AND
- 4. The requested dose is within FDA labeled dosing or supported in AASLD/IDSA dosing for the requested indication AND
- 5. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
  - C. Patient has an FDA labeled contraindication or hypersensitivity to the preferred agent: Mavyret for supported genotypes OR
  - D. Prescriber has provided information based on FDA approved labeling or AASLD/IDSA guidelines supporting the use of the non-preferred agent for the patient's diagnosis and genotype over the preferred agent: Mavyret for supported genotypes

## Age Restriction:

# **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, hepatologist or infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Duration of therapy: Based on FDA approved labeling or AASLD/IDSA guideline supported

Hetlioz LQ Suspension PA

# Drug Name(s)

Hetlioz Lq

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of Smith-Magenis Syndrome (SMS) confirmed by the presence of ONE of the following genetic mutations:
  - A. A heterozygous deletion of 17p11.2 OR
  - B. A heterozygous pathogenic variant involving RAI1 AND
- 2. The requested agent is being used to treat nighttime sleep disturbances associated with SMS

# Age Restriction:

Patient is 3 to 15 years of age

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist, sleep specialist, psychiatrist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

High Risk Medication PA - All Starts

# Drug Name(s)

Ascomp/Codeine

Benztropine Mesylate

Bonjesta

Butalbital/Acetaminophen/Caffeine/Codeine

Butalbital/Aspirin/Caffeine/Codeine

Carbinoxamine Maleate

Clemastine Fumarate

Cyproheptadine Hcl

Diclegis

Dicyclomine Hcl

Diphenoxylate/Atropine

Doxylamine Succinate/Pyridoxine Hcl

Fioricet/Codeine

Hydroxyzine Hcl

Hydroxyzine Pamoate

Lomotil

Promethazine Hcl Plain

Promethazine Hcl

Promethazine Vc

Promethazine/Phenylephrine

Promethegan

Ryvent

Scopolamine

Transderm-Scop

Trihexyphenidyl Hcl

Vistaril

### Indications:

All Medically-Accepted Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

# **Required Medical Information:**

PA does NOT apply to patients less than 65 years of age.

Criteria for approval require ALL of the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested high-risk medication AND

- 2. Prescriber has indicated that the benefits of the requested high-risk medication outweigh the risks for the patient AND
- 3. Prescriber has indicated that the risks and potential side effects of the requested high-risk medication have been discussed with the patient

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

HoFH PA - Juxtapid

#### Drug Name(s)

Juxtapid

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of homozygous familial hypercholesterolemia (HoFH) AND
- 2. Patient's diagnosis was confirmed by ONE of the following:
  - A. Genetic confirmation of bi-allelic pathogenic/likely pathogenic variants on different chromosomes at the LDLR, Apo-B, PCSK9, or LDLRAP1 genes or greater than or equal to 2 such variants at different loci OR
  - B. History of untreated LDL-C greater than 400 mg/dL (greater than 10 mmol/L) AND ONE of the following:
    - i. Patient has cutaneous or tendon xanthomas before the age of 10 years OR
    - ii. Untreated elevated LDL-C levels consistent with heterozygous familial hypercholesterolemia (HeFH) in both parents (or in digenic form, one parent may have normal LDL-C levels and the other may have LDL-C levels consistent with HoFH) AND
- 3. ONE of the following:
  - A. Patient is currently being treated with a lipid-lowering regimen in the last 90 days (i.e., rosuvastatin in combination with ezetimibe OR atorvastatin in combination with ezetimibe) OR
  - B. Patient has an intolerance or hypersensitivity to a lipid-lowering regimen (i.e., rosuvastatin in combination with ezetimibe OR atorvastatin in combination with ezetimibe) OR
  - C. Patient has an FDA labeled contraindication to a lipid-lowering regimen (i.e., rosuvastatin in combination with ezetimibe OR atorvastatin in combination with ezetimibe)

## Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cardiologist, endocrinologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

### Other Criteria:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of homozygous familial hypercholesterolemia (HoFH) AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. ONE of the following:
  - A. Patient is currently being treated with a lipid-lowering regimen in the last 90 days (i.e., rosuvastatin in combination with ezetimibe) OR

- B. Patient has an intolerance or hypersensitivity to a lipid-lowering regimen (i.e., rosuvastatin in combination with ezetimibe OR atorvastatin in combination with ezetimibe) OR
- C. Patient has an FDA labeled contraindication to a lipid-lowering regimen (i.e., rosuvastatin in combination with ezetimibe OR atorvastatin in combination with ezetimibe)

Horizant PA

# Drug Name(s)

Horizant

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. Patient has a diagnosis of Postherpetic Neuralgia (PHN) AND
    - ii. ONE of the following:
      - 1. Patient has tried and had an inadequate response to immediate-release gabapentin OR
      - 2. Patient has an intolerance or hypersensitivity to immediate-release gabapentin OR
      - 3. Patient has an FDA labeled contraindication to immediate-release gabapentin that is not expected to occur with the requested agent OR
  - B. BOTH of the following:
    - i. Patient has a diagnosis of moderate-to-severe primary Restless Legs Syndrome (RLS) AND
    - ii. BOTH of the following:
      - 1. ONE of the following:
        - a. Patient has tried and had an inadequate response to immediaterelease ropinirole OR
        - b. Patient has an intolerance or hypersensitivity to immediate-release ropinirole OR
        - c. Patient has an FDA labeled contraindication to immediate-release ropinirole AND
      - 2. ONE of the following:
        - a. Patient has tried and had an inadequate response to immediate-release pramipexole OR
        - b. Patient has an intolerance or hypersensitivity to immediate-release pramipexole OR
        - c. Patient has an FDA labeled contraindication to immediate-release pramipexole

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of Postherpetic Neuralgia (PHN) OR

- B. Patient has a diagnosis of moderate-to-severe primary Restless Legs Syndrome (RLS) AND
- 3. Patient has had clinical benefit with the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

**Hyftor PA** 

#### Drug Name(s)

Hyftor

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of facial angiofibroma associated with tuberous sclerosis complex (TSC) AND
- 2. Patient's diagnosis has been confirmed (i.e., medical records) by ONE of the following:
  - A. Patient has a pathogenic variant in the TSC1 gene or TSC2 gene confirmed by genetic testing OR
  - B. Patient has TWO of any of the following major features of TSC clinical diagnostic criteria:
    - i. hypomelanotic macules (greater than or equal to 3, at least 5 mm diameter)
    - ii. angiofibroma (greater than or equal to 3) or fibrous cephalic plaque
    - iii. ungual fibromas (greater than or equal to 2)
    - iv. shagreen patch
    - v. multiple retinal hamartomas
    - vi. multiple cortical tubers and/or radial migration lines
    - vii. subependymal nodule (greater than or equal to 2)
    - viii. subependymal giant cell astrocytoma
    - ix. cardiac rhabdomyoma
    - x. lymphangiomyomatosis (LAM)\*
    - xi. angiomyolipomas (greater than or equal to 2)\* OR
  - C. BOTH of the following:
    - i. Patient has ONE of any of the following major features of TSC clinical diagnostic criteria:
      - 1. hypomelanotic macules (greater than or equal to 3, at least 5 mm diameter)
      - 2. angiofibroma (greater than or equal to 3) or fibrous cephalic plaque
      - 3. ungual fibromas (greater than or equal to 2)
      - 4. shagreen patch
      - 5. multiple retinal hamartomas
      - 6. multiple cortical tubers and/or radial migration lines
      - 7. subependymal nodule (greater than or equal to 2)
      - 8. subependymal giant cell astrocytoma
      - 9. cardiac rhabdomyoma
      - 10. lymphangiomyomatosis (LAM)
      - 11. angiomyolipomas (greater than or equal to 2) AND

## Note:

\*A combination of the 2 major clinical features LAM and angiomyolipomas without other features does not meet criteria for a definite diagnosis.

Initial criteria continues: see Other Criteria

#### Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., dermatologist, geneticist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be 12 weeks for initial, 12 months for renewal

### Other Criteria:

- ii. Patient has TWO of any of the following minor features of TSC clinical diagnostic criteria:
  - 1. "confetti" skin lesions
  - 2. dental enamel pits (greater than or equal to 3)
  - 3. intraoral fibromas (greater than or equal to 2)
  - 4. retinal achromic patch
  - 5. multiple renal cysts
  - 6. nonrenal hamartomas
  - 7. sclerotic bone lesions AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of facial angiofibroma associated with tuberous sclerosis complex (TSC) AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Imiquimod PA

# Drug Name(s)

Imiquimod

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Actinic keratosis OR
  - B. Superficial basal cell carcinoma OR
  - C. External genital and/or perianal warts/condyloma acuminata OR
  - D. Squamous cell carcinoma OR
  - E. Basal cell carcinoma OR
  - F. Another indication that is supported in CMS approved compendia for the requested agent

## **Age Restriction:**

# **Prescriber Restrictions:**

# **Coverage Duration:**

4 months for Actinic keratosis, other diagnoses - see Other Criteria

### Other Criteria:

2 months for Superficial basal cell carcinoma, Squamous cell carcinoma, or Basal cell carcinoma

4 months for External genital and/or perianal warts/condyloma acuminata

12 months for All other diagnoses

Inbrija PA

# Drug Name(s)

Inbrija

### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. The requested agent will be used for intermittent treatment of OFF episodes in patients with Parkinson's disease AND
- 2. The requested agent will be used in combination with carbidopa/levodopa

# **Age Restriction:**

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Ingrezza PA

# Drug Name(s)

Ingrezza

**Indications:** 

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chorea associated with Huntington's disease AND BOTH of the following:
    - i. ONE of the following:
      - 1. Patient does NOT have a current diagnosis of depression OR
      - 2. Patient has a current diagnosis of depression and is being treated for depression AND
    - ii. ONE of the following:
      - 1. Patient does NOT have a diagnosis of passive suicidal ideation and/or behavior OR
      - 2. Patient has a diagnosis of passive suicidal ideation and/or behavior and must NOT be actively suicidal OR
  - B. Patient has a diagnosis of tardive dyskinesia AND ONE of the following:
    - i. Prescriber has reduced the dose of or discontinued any medications known to cause tardive dyskinesia (i.e., dopamine receptor blocking agents) OR
    - ii. Prescriber has provided clinical rationale indicating that a reduced dose or discontinuation of any medications known to cause tardive dyskinesia is not appropriate

# Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Injectable Oncology PA

## Drug Name(s)

Kanjinti

Mvasi

Ontruzant

Trazimera

Zirabev

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR C. ALL of the following:
    - i. Genetic testing has been completed, if required, for therapy with the requested agent and results indicate the requested agent is appropriate AND
    - ii. ONE of the following:
      - a. The requested agent is FDA labeled or supported by CMS approved compendia as first-line therapy for the requested indication OR
      - b. Patient has tried appropriate FDA labeled or CMS approved compendia supported therapy that are indicated as first-line therapy for the requested indication OR
      - c. Patient has an intolerance or hypersensitivity to the first-line therapy for the requested indication OR
      - d. Patient has an FDA labeled contraindication to the first-line therapy for the requested indication AND
    - iii. Patient does NOT have any FDA labeled contraindications to the requested agent AND
    - iv. Patient does NOT have any FDA labeled limitations of use that is not otherwise supported in NCCN guidelines

May also be subject to Part B versus Part D review.

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Insulin Pen Needle PA

# Drug Name(s)

Insulin Pen Needle

### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

This program will be implemented as a dynamic PA.

Criteria for approval require BOTH of the following:

- 1. The requested medical supply product will be used in the delivery of insulin to the body AND
- 2. Patient's medication history includes use of insulin within the past 180 days

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Insulin Syringe\_Needle PA

# Drug Name(s)

Insulin Syringe/Needle

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

This program will be implemented as a dynamic PA.

Criteria for approval require BOTH of the following:

- 1. The requested medical supply product will be used in the delivery of insulin to the body AND
- 2. Patient's medication history includes use of insulin within the past 180 days

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Intrarosa PA

# Drug Name(s)

Intrarosa

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Iron Chelating Agents PA - Exjade

# Drug Name(s)

Deferasirox (Exjade)

Exjade

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic iron overload due to a non-transfusion dependent thalassemia syndrome AND ONE of the following:
    - i. A liver iron (Fe) concentration (LIC) of at least 5 mg Fe per gram of dry weight OR
    - ii. A serum ferritin greater than 300 mcg/L OR
    - iii. MRI confirmation of iron deposition OR
  - B. Patient has a diagnosis of chronic iron overload due to blood transfusions AND
- 2. Patient will NOT be using the requested agent in combination with another iron chelating agent (e.g., deferiprone) for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of chronic iron overload due to a non-transfusion dependent thalassemia syndrome OR
  - B. Patient has a diagnosis of chronic iron overload due to blood transfusions AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another iron chelating agent (e.g., deferiprone) for the requested indication

# Age Restriction:

Patient is within the FDA labeled age for the requested agent for the requested indication

### **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

Iron Chelating Agents PA - Jadenu

# Drug Name(s)

Deferasirox (Jadenu)

Jadenu

Jadenu Sprinkle

### **Indications:**

All FDA-Approved Indications.

## Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic iron overload due to a non-transfusion dependent thalassemia syndrome AND ONE of the following:
    - i. A liver iron (Fe) concentration (LIC) of at least 5 mg Fe per gram of dry weight OR
    - ii. A serum ferritin greater than 300 mcg/L OR
    - iii. MRI confirmation of iron deposition OR
  - B. Patient has a diagnosis of chronic iron overload due to blood transfusions AND
- 2. Patient will NOT be using the requested agent in combination with another iron chelating agent (e.g., deferiprone) for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of chronic iron overload due to a non-transfusion dependent thalassemia syndrome OR
  - B. Patient has a diagnosis of chronic iron overload due to blood transfusions AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another iron chelating agent (e.g., deferiprone) for the requested indication

### Age Restriction:

Patient is within the FDA labeled age for the requested agent for the requested indication

## **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

Isturisa PA

## Drug Name(s)

Isturisa

#### **Indications:**

All FDA-Approved Indications.

#### **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of Cushing's disease AND
- 2. ONE of the following:
  - A. Patient had an inadequate response to pituitary surgical resection OR
  - B. Patient is NOT a candidate for pituitary surgical resection AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to pasireotide OR
  - B. Patient has an intolerance or hypersensitivity to pasireotide OR
  - C. Patient has an FDA labeled contraindication to pasireotide

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Cushing's disease AND
- 3. Patient has had clinical benefit with the requested agent

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Ivermectin Cream PA

# Drug Name(s)

Ivermectin Cream

Soolantra

### **Indications:**

All Medically-Accepted Indications.

# Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Ivermectin Tablet PA

# Drug Name(s)

Ivermectin Tablet

Stromectol

### **Indications:**

All Medically-Accepted Indications.

# Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

**Age Restriction:** 

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 4 months

Jatenzo PA

## Drug Name(s)

Jatenzo

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient's sex is male with a diagnosis of primary or secondary (hypogonadotropic) hypogonadism AND
- 2. ONE of the following:
  - A. Patient is NOT currently receiving testosterone replacement therapy AND has ONE of the following pretreatment levels:
    - i. Total serum testosterone level that is below the testing laboratory's lower limit of the normal range or is less than 300 ng/dL OR
    - ii. Free serum testosterone level that is below the testing laboratory's lower limit of the normal range OR
  - B. Patient is currently receiving testosterone replacement therapy AND has ONE of the following current levels:
    - i. Total serum testosterone level that is within the testing laboratory's normal range OR below the testing laboratory's lower limit of the normal range OR is less than 300  $\,\mathrm{ng/dL}$  OR
    - ii. Free serum testosterone level is within the testing laboratory's normal range OR below the testing laboratory's normal range AND
- 3. ONE of the following:
  - A. Patient will NOT be using the requested agent in combination with another androgen or anabolic steroid OR
  - B. Prescriber has provided information in support of therapy with more than one agent

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Joenja PA

# Drug Name(s)

Joenja

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of activated phosphoinositide 3-kinase (PI3K) delta syndrome (APDS) AND
- 2. Patient has a variant in either PIK3CD gene or PIK3R1 gene

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of activated phosphoinositide 3-kinase (PI3K) delta syndrome (APDS) AND
- 3. Patient has had clinical benefit with the requested agent

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., geneticist, immunologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Jornay PM PA

Drug Name(s)

Jornay Pm

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Jynarque PA

# Drug Name(s)

Jynarque

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of autosomal dominant polycystic kidney disease (ADPKD) confirmed by ONE of the following:
  - A. Ultrasound OR
  - B. MRI or CT scan OR
  - C. Genetic testing AND
- 2. Patient is at risk of rapid disease progression AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of autosomal dominant polycystic kidney disease (ADPKD) AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., nephrologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Kalydeco PA

# Drug Name(s)

Kalydeco

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of cystic fibrosis AND
- 2. ONE of the following:
  - A. Patient has ONE of the CFTR gene mutations or a mutation in the CFTR gene that is responsive based on in vitro data, as indicated in the FDA label, confirmed by genetic testing OR
  - B. Patient has another CFTR gene mutation(s) that is responsive to the requested agent, as indicated in the FDA label, confirmed by genetic testing AND
- 3. Patient is NOT homozygous for the F508del mutation AND
- 4. Patient will NOT be using the requested agent in combination with another CFTR modulator agent for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of cystic fibrosis AND
- 3. Patient has had improvement or stabilization with the requested agent [e.g., improvement in FEV1 from baseline, increase in weight/BMI, improvement from baseline Cystic Fibrosis Questionnaire-Revised (CFQ-R) Respiratory Domain score, improvements in respiratory symptoms (cough, sputum production, and difficulty breathing), and/or reduced number of pulmonary exacerbations] AND
- 4. Patient will NOT be using the requested agent in combination with another CFTR modulator agent for the requested indication

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cystic fibrosis, pulmonologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Kerendia PA

Drug Name(s)

Kerendia

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Leuprolide PA

# Drug Name(s)

Eligard

Leuprolide Acetate

Lupron Depot (1-Month)

Lupron Depot (3-Month)

Lupron Depot (4-Month)

Lupron Depot (6-Month)

Lupron Depot-Ped (1-Month)

Lupron Depot-Ped (3-Month)

Lupron Depot-Ped (6-Month)

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. BOTH of the following:
    - i. Patient is NOT currently being treated with the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent AND
- 3. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

**Age Restriction:** 

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Lidocaine Topical PA - Lidocaine Ointment

Drug Name(s)

Lidocaine Ointment

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Lidocaine Topical PA - Lidocaine Patch

Drug Name(s)

Lidocaine Patch

Lidoderm

Lidocan

Tridacaine II

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Lidocaine Topical PA - Lidocaine Solution

# Drug Name(s)

Lidocaine Solution

#### Indications:

All Medically-Accepted Indications.

# Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

- 1. The requested agent will be used for ONE of the following:
  - A. Topical anesthesia of accessible mucous membranes of the oral and nasal cavities OR
  - B. Topical anesthesia of accessible mucous membranes of proximal portions of the digestive tract OR
  - C. Another indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Lidocaine Topical PA - Lidocaine/prilocaine Cream

# Drug Name(s)

Lidocaine/Prilocaine

### **Indications:**

All Medically-Accepted Indications.

# Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

- 1. The requested agent will be used for ONE of the following:
  - A. Local analgesia on normal intact skin OR
  - B. Topical anesthetic for dermal procedures OR
  - C. Adjunctive anesthesia prior to local anesthetic infiltration in adult male genital skin OR
  - D. Anesthesia for minor procedures on female external genitalia OR
  - E. Another indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 months

Lidocaine Topical PA – Pliaglis

Drug Name(s)

Pliaglis

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. The requested agent will be used on intact skin of an adult patient to provide local analgesia for a superficial dermatological procedure (e.g., dermal filler injection, pulsed dye laser therapy, facial laser resurfacing, laser-assisted tattoo removal)

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Lidocaine Topical PA – ZTlido

Drug Name(s)

Ztlido

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Linezolid PA

## Drug Name(s)

Linezolid

Zyvox

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND ONE of the following:
  - a. The requested agent is prescribed by an infectious disease specialist or the prescriber has consulted with an infectious disease specialist on treatment of this patient OR
  - b. Patient has a documented infection due to vancomycin-resistant Enterococcus faecium OR
  - c. Patient has a diagnosis of pneumonia caused by Staphylococcus aureus or Streptococcus pneumoniae AND ONE of the following:
    - i. Patient has a documented infection that is resistant to TWO of the following: betalactams, macrolides, clindamycin, tetracyclines, or co-trimoxazole, OR that is resistant to vancomycin OR
    - ii. Patient has an intolerance or hypersensitivity to TWO of the following: beta-lactams, macrolides, clindamycin, tetracyclines, or co-trimoxazole OR
    - iii. Patient has an FDA labeled contraindication to TWO of the following: beta-lactams, macrolides, clindamycin, tetracyclines, or co-trimoxazole OR
    - iv. Patient has an intolerance or hypersensitivity to vancomycin OR
    - v. Patient has an FDA labeled contraindication to vancomycin OR
  - d. Patient has a documented skin and skin structure infection, including diabetic foot infections, caused by Staphylococcus aureus, Streptococcus pyogenes, or Streptococcus agalactiae AND ONE of the following:
    - i. Patient has a documented infection that is resistant to TWO of the following: betalactams, macrolides, clindamycin, tetracyclines, or co-trimoxazole, OR that is resistant to vancomycin at the site of infection OR
    - ii. Patient has an intolerance or hypersensitivity to TWO of the following: beta-lactams, macrolides, clindamycin, tetracyclines, or co-trimoxazole OR
    - iii. Patient has an FDA labeled contraindication to TWO of the following: beta-lactams, macrolides, clindamycin, tetracyclines, or co-trimoxazole OR

Criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 3 months

- iv. Patient has an intolerance or hypersensitivity to vancomycin OR
- v. Patient has an FDA labeled contraindication to vancomycin AND
- 2. Patient will NOT be using the requested agent in combination with Sivextro (tedizolid) for the same infection AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Livmarli PA

# Drug Name(s)

Livmarli

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of progressive familial intrahepatic cholestasis (PFIC) OR
  - B. Patient has a diagnosis of Alagille Syndrome (ALGS) AND
- 2. The requested agent will be used to treat cholestatic pruritus AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, hepatologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Lumryz PA

# Drug Name(s)

Lumryz

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of narcolepsy with cataplexy OR
  - B. Patient has a diagnosis of narcolepsy with excessive daytime sleepiness AND BOTH of the following:
    - i. ONE of the following:
      - a. Patient has tried and had an inadequate response to modafinil or armodafinil OR
      - b. Patient has an intolerance or hypersensitivity to modafinil or armodafinil OR
      - c. Patient has an FDA labeled contraindication to modafinil or armodafinil AND
    - ii. ONE of the following:
      - a. Patient has tried and had an inadequate response to ONE standard stimulant agent (e.g., methylphenidate) OR
      - b. Patient has an intolerance or hypersensitivity to ONE standard stimulant agent (e.g., methylphenidate) OR
      - c. Patient has an FDA labeled contraindication to ONE standard stimulant agent (e.g., methylphenidate) OR
  - C. Patient has another indication that is supported in CMS approved compendia for the requested agent

## Age Restriction:

Patient is 18 years of age or over

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Lupkynis PA

Drug Name(s)

Lupkynis

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of active lupus nephritis (LN) AND
- 2. Patient will continue standard LN therapy [corticosteroids (e.g., methylprednisolone, prednisone), immunosuppressives (e.g., azathioprine, mycophenolate)] in combination with the requested agent AND
- 3. Patient will NOT be using the requested agent in combination with cyclophosphamide

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of active lupus nephritis (LN) AND
- 3. Patient will continue standard LN therapy [corticosteroids (e.g., methylprednisolone, prednisone), immunosuppressives (e.g., azathioprine, mycophenolate)] in combination with the requested agent AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with cyclophosphamide

#### Age Restriction:

Patient is 18 years of age or over

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Mavyret PA

# Drug Name(s)

Mavyret

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of hepatitis C confirmed by serological markers OR
  - B. Patient is a hepatitis C virus (HCV) uninfected solid organ transplant recipient AND BOTH of the following:
    - i. Patient received an HCV viremic donor organ AND
    - ii. The requested agent is being used for prophylaxis AND
- 2. Prescriber has screened the patient for current or prior hepatitis B viral (HBV) infection and if positive, will monitor the patient for HBV flare-up or reactivation during and after treatment with the requested agent AND
- 3. The requested agent will be used in a treatment regimen and length of therapy that is supported in FDA approved labeling or AASLD/IDSA guidelines for the patient's diagnosis and genotype AND
- 4. The requested dose is within FDA labeled dosing or supported in AASLD/IDSA guideline dosing for the requested indication

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, hepatologist or infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Duration of therapy: Based on FDA approved labeling or AASLD/IDSA guideline supported

Memantine ER PA

# Drug Name(s)

Memantine Hcl Er

Namenda Xr

### **Indications:**

All Medically-Accepted Indications.

# Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

PA does NOT apply to patients greater than or equal to 30 years of age Criteria for approval require the following:

- 1. Patient is younger than 30 years of age AND ONE of the following:
  - A. Patient has a diagnosis of moderate to severe dementia of the Alzheimer's type OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Memantine PA

# Drug Name(s)

Memantine Hcl Titration Pak

Memantine Hcl

Namenda

Namenda Titration Pak

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

PA does NOT apply to patients greater than or equal to 30 years of age Criteria for approval require the following:

- 1. Patient is younger than 30 years of age AND ONE of the following:
  - A. Patient has a diagnosis of moderate to severe dementia of the Alzheimer's type OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Methamphetamine PA

# Drug Name(s)

Methamphetamine Hcl

# **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

# **Exclusion Criteria:**

Requested agent will be used to promote weight loss AND FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Methylin PA

# Drug Name(s)

Methylin

Methylphenidate Hcl (Methylin)

**Indications:** 

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Methylphenidate Capsule PA

Drug Name(s)

Metadate Cd

Methylphenidate Hcl Cd

Methylphenidate Hcl Er Capsule

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Methylphenidate Chewable PA

Drug Name(s)

Methylphenidate Hcl Chewable

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Methylphenidate ER Tablet PA

# Drug Name(s)

Methylphenidate Hcl Er Tablet

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Methylphenidate LA Capsule PA

# Drug Name(s)

Methylphenidate Hcl Er (La)

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Methylphenidate Patch PA

Drug Name(s)

Daytrana

MethylphenidatePatch

**Indications:** 

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Miebo PA

# Drug Name(s)

Miebo

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Mifepristone PA

# Drug Name(s)

Korlym

Mifepristone

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of Cushing's syndrome AND
- 2. ONE of the following:
  - A. Patient has type 2 diabetes mellitus OR
  - B. Patient has glucose intolerance as defined by a 2-hour glucose tolerance test plasma glucose value of 140-199 mg/dL AND
- 3. ONE of the following:
  - A. Patient had an inadequate response to surgical resection OR
  - B. Patient is NOT a candidate for surgical resection

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Cushing's syndrome AND
- 3. Patient has had clinical benefit with the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

## **Coverage Duration:**

Approval will be for 12 months

Migranal PA

### Drug Name(s)

Dihydroergotamine Mesylate Spray

Migranal

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. The requested agent will be used for the treatment of acute migraine with or without aura AND
- 2. ONE of the following:
  - A. Patient has tried and had an inadequate response to TWO triptan agents with differing active ingredients (e.g., sumatriptan, rizatriptan) OR
  - B. Patient has an intolerance or hypersensitivity to TWO triptan agents with differing active ingredients OR
  - C. Patient has an FDA labeled contraindication to TWO triptan agents with differing active ingredients AND
- 3. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, acute CGRP)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. The requested agent will be used for the treatment of acute migraine with or without aura AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, acute CGRP)

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Modafinil PA

# Drug Name(s)

Modafinil

Provigil

# **Indications:**

All Medically-Accepted Indications.

# Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another target agent (i.e., armodafinil)

# Age Restriction:

Patient is 17 years of age or over

# **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

Mounjaro PA

# Drug Name(s)

Mounjaro

### **Indications:**

All FDA-Approved Indications.

### **Off-Label Uses:**

### **Exclusion Criteria:**

Requested agent will be used for weight loss alone

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of type 2 diabetes mellitus AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 180 days OR
  - C. ALL of the following:
    - i. ONE of the following:
      - 1. Patient's medication history includes use of a non glucagon-like peptide-1 (GLP-1) oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) within the past 90 days OR
      - 2. Patient had an ineffective treatment response to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 3. Patient has an intolerance or hypersensitivity to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 4. Patient has an FDA labeled contraindication to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent  $\ensuremath{\mathsf{AND}}$
    - iii. Patient will NOT be using the requested agent in combination with another GLP-1 agonist agent, or an agent containing a GLP-1 agonist AND
    - iv. Patient will NOT be using the requested agent in combination with an agent containing a dipeptidyl peptidase-4 (DPP-4) inhibitor

### Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

MS PA – Avonex

# Drug Name(s)

Avonex

Avonex Pen

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

MS PA – Bafiertam

# Drug Name(s)

Bafiertam

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 months

MS PA – Betaseron

## Drug Name(s)

Betaseron

Extavia

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

MS PA - Dimethyl Fumarate

# Drug Name(s)

Dimethyl Fumarate

Dimethyl Fumarate Starterpack

Tecfidera

Tecfidera Starter Pack

#### Indications:

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

MS PA - Fingolimod

# Drug Name(s)

Fingolimod

Gilenya

### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication AND
- 3. Prescriber has performed an electrocardiogram within 6 months prior to initiating treatment

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

MS PA – Glatiramer

# Drug Name(s)

Copaxone

Glatiramer Acetate

Glatopa

# **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

MS PA – Kesimpta

Drug Name(s)

Kesimpta

**Indications:** 

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

MS PA - Mavenclad

# Drug Name(s)

Mavenclad

**Indications:** 

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) for the requested indication AND
- 3. The requested dose is within FDA labeled dosing for the requested indication AND
- 4. The total cumulative duration of treatment with Mavenclad (cladribine) has not exceeded 4 treatment cycles

# Age Restriction:

#### **Prescriber Restrictions:**

# **Coverage Duration:**

No prior use approve 2 years, Prior use approve remainder of 2 years of total cumulative therapy **Other Criteria:** 

MS PA – Mayzent

# Drug Name(s)

Mayzent

Mayzent Starter Pack

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

MS PA – Plegridy

# Drug Name(s)

Plegridy

Plegridy Starter Pack

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

MS PA – Ponvory

# Drug Name(s)

Ponvory

Ponvory 14-Day Starter Pack

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

MS PA – Rebif

## Drug Name(s)

Rebif

Rebif Rebidose

Rebif Rebidose Titration Pack

**Rebif Titration Pack** 

#### Indications:

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

MS PA – Tascenso

### Drug Name(s)

Tascenso Odt

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication AND
- 3. Prescriber has performed an electrocardiogram within 6 months prior to initiating treatment

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

MS PA - Teriflunomide

# Drug Name(s)

Aubagio

Teriflunomide

### Indications:

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be for 12 months

MS PA – Vumerity

## Drug Name(s)

Vumerity

**Indications:** 

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

MS PA – Zeposia

## Drug Name(s)

Zeposia

Zeposia Starter Kit

Zeposia 7-Day Starter Pack

### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication AND
- 3. ONE of the following:
  - i. The requested diagnosis is NOT moderately to severely active ulcerative colitis (UC) OR
  - ii. The requested diagnosis is moderately to severely active UC AND BOTH of the following:
    - 1. ONE of the following:
      - a. Patient has tried and had an inadequate response to at least ONE conventional agent (e.g., 5-aminosalicylates [including balsalazide, mesalamine, olsalazine, sulfasalazine], mercaptopurine, azathioprine, corticosteroids [including budesonide EC capsule]) used in the treatment of UC OR
      - b. Patient has severely active UC OR
      - c. Patient has an intolerance or hypersensitivity to ONE of the conventional agents used in the treatment of UC OR
      - d. Patient has an FDA labeled contraindication to at least ONE of the conventional agents used in the treatment of UC AND
    - 2. ONE of the following:
      - a. Patient has tried and had an inadequate response to ONE preferred biologic agent (Hadlima or Stelara) for the treatment of UC OR
      - b. Patient has an intolerance or hypersensitivity to ONE preferred biologic agent (Hadlima or Stelara) OR
      - c. Patient has an FDA labeled contraindication to ONE preferred biologic agent (Hadlima or Stelara)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another disease modifying agent (DMA) or a biologic immunomodulator for the requested indication

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Mulpleta PA

# Drug Name(s)

Mulpleta

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of thrombocytopenia AND
- 2. Patient has chronic liver disease AND
- 3. Patient has a platelet count less than 50 X 10^9/L AND
- 4. Patient is scheduled to undergo a procedure with an associated risk of bleeding (e.g., gastrointestinal endoscopy, liver biopsy, bronchoscopy, dental procedure) AND
- 5. The requested dose is within FDA labeled dosing for the requested indication AND
- 6. The length of therapy of the requested agent is within the FDA labeled duration for the requested indication

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 1 month

Myalept PA

### Drug Name(s)

Myalept

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has leptin deficiency associated with a diagnosis of either congenital generalized lipodystrophy (CGL) or acquired generalized lipodystrophy (AGL) AND
- 2. Prescriber has provided the patient's baseline levels for HbA1C, triglycerides, and fasting insulin, measured prior to beginning therapy with the requested agent AND
- 3. Patient also has at least ONE of the complications related to lipodystrophy: diabetes mellitus, hypertriglyceridemia (200 mg/dL or higher), and/or high fasting insulin (30 microunits/mL or higher) AND
- 4. Patient has tried and had an inadequate response to maximum tolerable dosing of a conventional agent for the additional diagnosis AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has leptin deficiency associated with a diagnosis of either congenital generalized lipodystrophy (CGL) or acquired generalized lipodystrophy (AGL) AND
- 3. Patient has had improvement or stabilization with the requested agent as indicated by change from baseline level of at least ONE of the following:
  - A. HbA1C
  - B. Triglycerides
  - C. Fasting insulin AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., endocrinologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Conventional agent examples include:

Hypertriglyceridemia: statins, fenofibrates, Omega-3-Acid Ethyl Esters (generic Lovaza)

Diabetes/high fasting insulin: insulin, sulfonylurea/sulfonylurea combination, metformin/metformin combination

Myfembree PA

# Drug Name(s)

Myfembree

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. ONE of the following:
- A. BOTH of the following:
- i. Patient has a diagnosis of heavy menstrual bleeding associated with uterine leiomyomas (fibroids) AND
- ii. Patient's diagnosis of uterine fibroids was confirmed via imaging (e.g., ultrasound) OR
- B. Patient has a diagnosis of moderate to severe pain associated with endometriosis AND
- 2. Patient is premenopausal AND
- 3. Patient will NOT be using the requested agent in combination with another GnRH antagonist agent [e.g., Orilissa (elagolix), Oriahnn (elagolix, estradiol, norethindrone acetate)] for the requested indication AND
- 4. The requested dose is within FDA labeled dosing for the requested indication AND
- 5. ONE of the following:
- A. Patient is initiating therapy with the requested agent OR
- B. BOTH of the following:
- i. Patient is continuing therapy with the requested agent and the prescriber has provided information indicating the number of months the patient has been on therapy with the requested agent AND
- ii. The total duration of treatment with the requested agent has NOT exceeded 24 months per lifetime

### **Age Restriction:**

### **Prescriber Restrictions:**

# **Coverage Duration:**

For no prior Myfembree use:

24 months

For prior Myfembree use:

Remainder of 24 months

Nexletol PA

## Drug Name(s)

Nexletol

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of heterozygous familial hypercholesterolemia (HeFH) AND ONE of the following:
    - i. Genetic confirmation of one mutant allele at the LDLR, Apo-B, PCSK9, or 1/LDLRAP1 gene OR
    - ii. Pretreatment LDL-C greater than 190 mg/dL (greater than 4.9 mmol/L) OR
    - iii. Patient has clinical manifestations of HeFH (e.g., cutaneous xanthomas, tendon xanthomas, corneal arcus, tuberous xanthoma, or xanthelasma) OR
    - iv. Patient has "definite" or "possible" familial hypercholesterolemia as defined by the Simon Broome criteria OR
    - v. Patient has a Dutch Lipid Clinic Network criteria score of greater than 5 OR
  - B. The requested agent will be used to reduce the risk of myocardial infarction and coronary revascularization AND ONE of the following:
    - i. Patient has a diagnosis of established atherosclerotic cardiovascular disease (ASCVD) defined as having ONE of the following:
      - a. Acute coronary syndrome
      - b. History of myocardial infarction
      - c. Stable or unstable angina
      - d. Coronary or other arterial revascularization
      - e. Stroke
      - f. Transient ischemic attack
      - g. Peripheral arterial disease, including aortic aneurysm, presumed to be of atherosclerotic origin OR
    - ii. Patient has a diagnosis of high risk for a cardiovascular disease (CVD) event but without established CVD OR
  - C. Patient has a diagnosis of primary hyperlipidemia, [not associated with HeFH, established cardiovascular disease (CVD), or high risk for CVD event without established CVD] AND
- 2. ONE of the following:
  - A. Patient is on statin therapy OR
  - B. Patient has an intolerance to TWO different statins OR
  - C. Patient has an FDA labeled contraindication to a statin

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cardiologist, endocrinologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of heterozygous familial hypercholesterolemia (HeFH) OR
  - B. The requested agent will be used to reduce the risk of myocardial infarction and coronary revascularization AND ONE of the following:
    - i. Patient has a diagnosis of established atherosclerotic cardiovascular disease (ASCVD) OR
    - ii. Patient has a diagnosis of high risk for a cardiovascular disease (CVD) event but without established CVD OR
  - C. Patient has a diagnosis of primary hyperlipidemia, [not associated with HeFH, established cardiovascular disease (CVD), or high risk for CVD event without established CVD] AND
- 3. ONE of the following:
  - A. Patient is on statin therapy OR
  - B. Patient has an intolerance to TWO different statins OR
  - C. Patient has an FDA labeled contraindication to a statin AND
- 4. Patient has had clinical benefit with the requested agent

Nexlizet PA

## Drug Name(s)

Nexlizet

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of heterozygous familial hypercholesterolemia (HeFH) AND ONE of the following:
    - i. Genetic confirmation of one mutant allele at the LDLR, Apo-B, PCSK9, or 1/LDLRAP1 gene OR
    - ii. Pretreatment LDL-C greater than 190 mg/dL (greater than 4.9 mmol/L) OR
    - iii. Patient has clinical manifestations of HeFH (e.g., cutaneous xanthomas, tendon xanthomas, corneal arcus, tuberous xanthoma, or xanthelasma) OR
    - iv. Patient has "definite" or "possible" familial hypercholesterolemia as defined by the Simon Broome criteria OR
    - v. Patient has a Dutch Lipid Clinic Network criteria score of greater than 5 OR
  - B. The requested agent will be used to reduce the risk of myocardial infarction and coronary revascularization AND ONE of the following:
    - i. Patient has a diagnosis of established atherosclerotic cardiovascular disease (ASCVD) defined as having ONE of the following:
      - a. Acute coronary syndrome
      - b. History of myocardial infarction
      - c. Stable or unstable angina
      - d. Coronary or other arterial revascularization
      - e. Stroke
      - f. Transient ischemic attack
      - g. Peripheral arterial disease, including aortic aneurysm, presumed to be of atherosclerotic origin OR
    - ii. Patient has a diagnosis of high risk for a cardiovascular disease (CVD) event but without established CVD OR
  - C. Patient has a diagnosis of primary hyperlipidemia, [not associated with HeFH, established cardiovascular disease (CVD), or high risk for CVD event without established CVD] AND
- 2. ONE of the following:
- A. Patient is on statin therapy OR
- B. Patient has an intolerance to TWO different statins OR
- C. Patient has an FDA labeled contraindication to a statin

## Age Restriction:

**Prescriber Restrictions:** 

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cardiologist, endocrinologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of heterozygous familial hypercholesterolemia (HeFH) OR
  - B. The requested agent will be used to reduce the risk of myocardial infarction and coronary revascularization AND ONE of the following:
    - i. Patient has a diagnosis of established atherosclerotic cardiovascular disease (ASCVD) OR
    - ii. Patient has a diagnosis of high risk for a cardiovascular disease (CVD) event but without established CVD OR
  - C. Patient has a diagnosis of primary hyperlipidemia, [not associated with HeFH, established cardiovascular disease (CVD), or high risk for CVD event without established CVD] AND
- 3. ONE of the following:
  - A. Patient is on statin therapy OR
  - B. Patient has an intolerance to TWO different statins OR
  - C. Patient has an FDA labeled contraindication to a statin AND
- 4. Patient has had clinical benefit with the requested agent

Nocdurna PA

# Drug Name(s)

Nocdurna

Indications:

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of nocturia due to nocturnal polyuria (awakening at least two times per night to void) AND
- 2. Diagnosis was confirmed by a nighttime urine production greater than one third of 24-hour urine collection AND
- 3. Patient's serum sodium concentration is within normal range [between 135 to 145 mEq/L (mmol/L) or within testing laboratory's normal range]

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of nocturia due to nocturnal polyuria (awakening at least two times per night to void) AND
- 3. Patient's serum sodium concentration is within normal range [between 135 to 145 mEq/L (mmol/L) or within testing laboratory's normal range] AND
- 4. Patient has had clinical benefit with the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Nourianz PA

## Drug Name(s)

Nourianz

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. The requested agent will be used as adjunctive treatment in patients with Parkinson's disease experiencing "off" episodes AND
- 2. The requested agent will be used in combination with levodopa/carbidopa agents

## Age Restriction:

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Nucala PA

### Drug Name(s)

Nucala

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of severe asthma with an eosinophilic phenotype AND the following:
    - i. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LRTA, LAMA, theophylline) in combination with the requested agent OR
  - B. Patient has a diagnosis of eosinophilic granulomatosis with polyangiitis (EGPA) AND the following:
    - i. ONE of the following:
      - a. Patient is currently being treated with a maximally tolerated oral corticosteroid OR
      - b. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to an oral corticosteroid OR
  - C. Patient has a diagnosis of hypereosinophilic syndrome (HES) for 6 months or more without an identifiable non-hematologic secondary cause AND the following:
    - i. ONE of the following:
      - a. Patient is currently being treated with a maximally tolerated oral corticosteroid OR
      - b. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to an oral corticosteroid OR
  - D. Patient has a diagnosis of chronic rhinosinusitis with nasal polyps (CRSwNP) AND
- 2. Patient will NOT be using the requested agent in combination with Xolair, Dupixent, or with another injectable interleukin 5 (IL-5) inhibitor (e.g., Cinqair, Fasenra) for the requested indication AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

## **Age Restriction:**

For diagnosis of severe asthma with an eosinophilic phenotype, patient is 6 years of age or over. For diagnosis of EGPA, patient is 18 years of age or over. For diagnosis of HES, patient is 12 years of age or over. For diagnosis of CRSwNP, patient is 18 years of age or over.

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, immunologist, otolaryngologist, pathologist, pulmonologist, rheumatologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of severe asthma with an eosinophilic phenotype AND the following:
    - i. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LTRA, LAMA, theophylline) in combination with the requested agent OR
  - B. Patient has a diagnosis of eosinophilic granulomatosis with polyangiitis (EGPA) AND the following:
    - i. ONE of the following:
      - a. Patient is currently being treated with maintenance therapy with oral corticosteroid OR
      - b. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to oral corticosteroid OR
  - C. Patient has a diagnosis of hypereosinophilic syndrome (HES) for 6 months or more without an identifiable non-hematologic secondary cause AND the following:
    - i. ONE of the following:
      - a. Patient is currently being treated with maintenance therapy with oral corticosteroid OR
      - b. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to oral corticosteroid OR
  - D. Patient has a diagnosis of chronic rhinosinusitis with nasal polyps (CRSwNP) AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with Xolair, Dupixent, or with another injectable interleukin 5 (IL-5) inhibitor (e.g., Cinqair, Fasenra) for the requested indication AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

Nuedexta PA

# Drug Name(s)

Nuedexta

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of pseudobulbar affect OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Nuplazid PA

Drug Name(s)

Nuplazid

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Nurtec PA

### Drug Name(s)

Nurtec

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of migraine AND
- 2. ONE of the following:
  - A. The requested agent is being used for the treatment of acute migraine with or without aura AND BOTH of the following:
    - i. ONE of the following:
      - a. Patient has tried and had an inadequate response to a triptan (e.g., sumatriptan, rizatriptan) agent OR
      - b. Patient has an intolerance, or hypersensitivity to a triptan OR
      - c. Patient has an FDA labeled contraindication to a triptan AND
    - ii. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, ergotamine, acute CGRP) OR
  - B. The requested agent is being used for migraine prophylaxis AND BOTH of the following:
    - i. Patient has 4 or more migraine headache days per month AND
    - ii. Patient will NOT be using the requested agent in combination with another calcitonin gene-related peptide (CGRP) agent for migraine prophylaxis

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of migraine AND
- 3. ONE of the following:
  - A. The requested agent is being used for the treatment of acute migraine with or without aura AND BOTH of the following:
    - i. Patient has had clinical benefit with the requested agent AND
    - ii. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, ergotamine, acute CGRP) OR
  - B. The requested agent is being used for migraine prophylaxis AND BOTH of the following:
    - i. Patient has had clinical benefit with the requested agent AND
    - ii. Patient will NOT be using the requested agent in combination with another calcitonin gene-related peptide (CGRP) agent for migraine prophylaxis

### Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 Months

Ocaliva PA

## Drug Name(s)

Ocaliva

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of primary biliary cholangitis (PBC) confirmed by at least TWO of the following:
  - A. There is biochemical evidence of cholestasis with an alkaline phosphatase (ALP) elevation
  - B. Presence of antimitochondrial antibody (AMA): a titer greater than 1:80 OR a level that is above the testing laboratory's upper limit of the normal range
  - C. If the AMA is negative or present only in low titer (less than or equal to 1:80), presence of other PBC-specific autoantibodies, including sp100 or gp210
  - D. Histologic evidence of nonsuppurative destruction cholangitis and destruction of interlobular bile ducts AND
- 2. ONE of the following:
  - A. Patient does NOT have cirrhosis OR
  - B. Patient has compensated cirrhosis with NO evidence of portal hypertension AND
- 3. Prescriber has measured the patient's alkaline phosphatase (ALP) level AND total bilirubin level AND
- 4. ONE of the following:
  - A. BOTH of the following:
    - i. Patient has tried and had an inadequate response to ursodiol AND
    - ii. The requested agent will be used in combination with ursodiol OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to ursodiol

# Age Restriction:

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of primary biliary cholangitis (PBC) AND
- 3. ONE of the following:
  - A. Patient does NOT have cirrhosis OR
  - B. Patient has compensated cirrhosis with NO evidence of portal hypertension AND
- 4. ONE of the following:
  - A. The requested agent will be used in combination with ursodiol OR
  - B. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to ursodiol AND

- 5. Patient has had improvements or stabilization with the requested agent as indicated by BOTH of the following:
  - A. Decrease in alkaline phosphatase (ALP) level from baseline AND
  - B. Total bilirubin is less than or equal to the upper limit of normal (ULN)

Ofev PA

### Drug Name(s)

Ofev

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. Patient has a diagnosis of idiopathic pulmonary fibrosis (IPF) AND
    - ii. Patient has no known explanation for interstitial lung disease (ILD) or pulmonary fibrosis (e.g., radiation, drugs, metal dusts, sarcoidosis, or any connective tissue disease known to cause ILD) OR
  - B. BOTH of the following:
    - i. Patient has a diagnosis of systemic sclerosis-associated interstitial lung disease (SSc-ILD) AND
    - ii. Patient's diagnosis has been confirmed on high-resolution computed tomography (HRCT) or chest radiography scans OR
  - C. BOTH of the following:
    - i. Patient has a diagnosis of chronic fibrosing interstitial lung disease (ILD) with a progressive phenotype AND
    - ii. Patient's diagnosis has been confirmed on high-resolution computed tomography (HRCT)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of ONE of the following:
  - A. Idiopathic pulmonary fibrosis (IPF) OR
  - B. Systemic sclerosis-associated interstitial lung disease (SSc-ILD) OR
  - C. Chronic fibrosing interstitial lung disease (ILD) with a progressive phenotype AND
- 3. Patient has had clinical benefit with the requested agent

# Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., pathologist, pulmonologist, radiologist, rheumatologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Omnipod PA

# Drug Name(s)

Omnipod 5 Kit

Omnipod 5 Pods

Omnipod Classic Kit

Omnipod Classic Pods

Omnipod Dash Kit

Omnipod Dash Pods

Omnipod Go

## **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of diabetes mellitus AND
- 2. Patient is on an insulin regimen of 3 or more injections per day AND
- 3. ONE of the following:
  - A. Patient is testing glucose levels 4 or more times per day OR
  - B. Patient is using a continuous glucose monitor (CGM)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of diabetes mellitus AND
- 3. Patient has had clinical benefit with the requested agent (e.g., stable or improved glycemic control)

# Age Restriction:

#### **Prescriber Restrictions:**

## **Coverage Duration:**

Approval will be for 12 months

Ophthalmic Immunomodulators PA – Cequa

# Drug Name(s)

Cequa

## Indications:

All Medically-Accepted Indications.

## **Off-Label Uses:**

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Ophthalmic Immunomodulators PA – Verkazia

# Drug Name(s)

Verkazia

### Indications:

All Medically-Accepted Indications.

### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Ophthalmic Immunomodulators PA – Vevye

# Drug Name(s)

Vevye

# Indications:

All Medically-Accepted Indications.

### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Ophthalmic Immunomodulators PA – Xiidra

# Drug Name(s)

Xiidra

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Opioids ER PA - Buprenorphine Pain

# Drug Name(s)

Belbuca

Buprenorphine

**Butrans** 

### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic cancer-related pain OR
  - B. Patient has a diagnosis of pain due to sickle cell disease OR
  - C. Patient is undergoing treatment of chronic non-cancer pain AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
    - iii. ALL of the following:
      - a. Prescriber has completed a formal, consultative evaluation including BOTH of the following:
        - 1. Diagnosis AND
        - 2. A medical history which includes previous and current pharmacological and non-pharmacological therapy for the requested diagnosis AND
      - b. The requested agent is NOT prescribed as an as-needed (prn) analgesic AND
      - c. Prescriber has confirmed that a patient-specific pain management plan is on file for the patient AND
      - d. ONE of the following:
        - 1. Patient's medication history includes use of an immediate-acting opioid OR
        - 2. Patient has an intolerance or hypersensitivity to an immediate-acting opioid OR
        - 3. Patient has an FDA labeled contraindication to an immediate-acting opioid AND
      - e. Prescriber has reviewed the patient's records in the state's prescription drug monitoring program (PDMP) AND has determined that the opioid dosages and combinations of opioids and other controlled substances within the patient's records do NOT indicate the patient is at high risk for overdose AND
      - f. Patient does NOT have any FDA labeled contraindications to the requested agent

#### Age Restriction:

Prescriber Restrictions:
Coverage Duration:
Approval will be for 12 months

Opioids ER PA - Fentanyl Patch

### Drug Name(s)

Fentanyl

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic cancer-related pain OR
  - B. Patient has a diagnosis of pain due to sickle cell disease OR
  - C. Patient is undergoing treatment of chronic non-cancer pain AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past  $90\ \text{days}\ \text{OR}$
    - ii. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
    - iii. ALL of the following:
      - a. Prescriber has completed a formal, consultative evaluation including BOTH of the following:
        - 1. Diagnosis AND
        - 2. A medical history which includes previous and current pharmacological and non-pharmacological therapy for the requested diagnosis AND
      - b. The requested agent is NOT prescribed as an as-needed (prn) analgesic AND
      - c. Prescriber has confirmed that a patient-specific pain management plan is on file for the patient AND
      - d. ONE of the following:
        - 1. Patient's medication history includes use of an immediate-acting opioid OR
        - 2. Patient has an intolerance or hypersensitivity to an immediate-acting opioid OR
        - 3. Patient has an FDA labeled contraindication to an immediate-acting opioid AND
      - e. Prescriber has reviewed the patient's records in the state's prescription drug monitoring program (PDMP) AND has determined that the opioid dosages and combinations of opioids and other controlled substances within the patient's records do NOT indicate the patient is at high risk for overdose AND
      - f. Patient does NOT have any FDA labeled contraindications to the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

Opioids ER PA - Hydrocodone

### Drug Name(s)

Hydrocodone Bitartrate Er

Hysingla Er

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic cancer-related pain OR
  - B. Patient has a diagnosis of pain due to sickle cell disease OR
  - C. Patient is undergoing treatment of chronic non-cancer pain AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
    - iii. ALL of the following:
      - a. Prescriber has completed a formal, consultative evaluation including BOTH of the following:
        - 1. Diagnosis AND
        - 2. A medical history which includes previous and current pharmacological and non-pharmacological therapy for the requested diagnosis AND
      - b. The requested agent is NOT prescribed as an as-needed (prn) analgesic AND
      - c. Prescriber has confirmed that a patient-specific pain management plan is on file for the patient AND
      - d. ONE of the following:
        - 1. Patient's medication history includes use of an immediate-acting opioid OR
        - 2. Patient has an intolerance or hypersensitivity to an immediate-acting opioid OR
        - 3. Patient has an FDA labeled contraindication to an immediate-acting opioid AND
      - e. Prescriber has reviewed the patient's records in the state's prescription drug monitoring program (PDMP) AND has determined that the opioid dosages and combinations of opioids and other controlled substances within the patient's records do NOT indicate the patient is at high risk for overdose AND
      - f. Patient does NOT have any FDA labeled contraindications to the requested agent

### Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Opioids ER PA - Hydromorphone

### Drug Name(s)

Hydromorphone Hcl Er

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic cancer-related pain OR
  - B. Patient has a diagnosis of pain due to sickle cell disease OR
  - C. Patient is undergoing treatment of chronic non-cancer pain AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
    - iii. ALL of the following:
      - a. Prescriber has completed a formal, consultative evaluation including BOTH of the following:
        - 1. Diagnosis AND
        - 2. A medical history which includes previous and current pharmacological and non-pharmacological therapy for the requested diagnosis AND
      - b. The requested agent is NOT prescribed as an as-needed (prn) analgesic AND
      - c. Prescriber has confirmed that a patient-specific pain management plan is on file for the patient AND
      - d. ONE of the following:
        - 1. Patient's medication history includes use of an immediate-acting opioid OR
        - 2. Patient has an intolerance or hypersensitivity to an immediate-acting opioid OR
        - 3. Patient has an FDA labeled contraindication to an immediate-acting opioid AND
      - e. Prescriber has reviewed the patient's records in the state's prescription drug monitoring program (PDMP) AND has determined that the opioid dosages and combinations of opioids and other controlled substances within the patient's records do NOT indicate the patient is at high risk for overdose AND
      - f. Patient does NOT have any FDA labeled contraindications to the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

Opioids ER PA - Morphine

### Drug Name(s)

Morphine Sulfate Er

Ms Contin

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic cancer-related pain OR
  - B. Patient has a diagnosis of pain due to sickle cell disease OR
  - C. Patient is undergoing treatment of chronic non-cancer pain AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
    - iii. ALL of the following:
      - a. Prescriber has completed a formal, consultative evaluation including BOTH of the following:
        - 1. Diagnosis AND
        - 2. A medical history which includes previous and current pharmacological and non-pharmacological therapy for the requested diagnosis AND
      - b. The requested agent is NOT prescribed as an as-needed (prn) analgesic AND
      - c. Prescriber has confirmed that a patient-specific pain management plan is on file for the patient AND
      - d. ONE of the following:
        - 1. Patient's medication history includes use of an immediate-acting opioid OR
        - 2. Patient has an intolerance or hypersensitivity to an immediate-acting opioid OR
        - 3. Patient has an FDA labeled contraindication to an immediate-acting opioid AND
      - e. Prescriber has reviewed the patient's records in the state's prescription drug monitoring program (PDMP) AND has determined that the opioid dosages and combinations of opioids and other controlled substances within the patient's records do NOT indicate the patient is at high risk for overdose AND
      - f. Patient does NOT have any FDA labeled contraindications to the requested agent

### Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Opioids ER PA - Oxycodone

### Drug Name(s)

Oxycodone Hcl Er

Oxycontin

Xtampza Er

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic cancer-related pain OR
  - B. Patient has a diagnosis of pain due to sickle cell disease OR
  - C. Patient is undergoing treatment of chronic non-cancer pain AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
    - iii. ALL of the following:
      - a. Prescriber has completed a formal, consultative evaluation including BOTH of the following:
        - 1. Diagnosis AND
        - 2. A medical history which includes previous and current pharmacological and non-pharmacological therapy for the requested diagnosis AND
      - b. The requested agent is NOT prescribed as an as-needed (prn) analgesic AND
      - c. Prescriber has confirmed that a patient-specific pain management plan is on file for the patient AND
      - d. ONE of the following:
        - 1. Patient's medication history includes use of an immediate-acting opioid OR
        - 2. Patient has an intolerance or hypersensitivity to an immediate-acting opioid OR
        - 3. Patient has an FDA labeled contraindication to an immediate-acting opioid AND
      - e. Prescriber has reviewed the patient's records in the state's prescription drug monitoring program (PDMP) AND has determined that the opioid dosages and combinations of opioids and other controlled substances within the patient's records do NOT indicate the patient is at high risk for overdose AND
      - f. Patient does NOT have any FDA labeled contraindications to the requested agent

#### Age Restriction:

Prescriber Restrictions:
Coverage Duration:
Approval will be for 12 months

Opioids ER PA - Oxymorphone

### Drug Name(s)

Oxymorphone Hcl Er

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic cancer-related pain OR
  - B. Patient has a diagnosis of pain due to sickle cell disease OR
  - C. Patient is undergoing treatment of chronic non-cancer pain AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
    - iii. ALL of the following:
      - a. Prescriber has completed a formal, consultative evaluation including BOTH of the following:
        - 1. Diagnosis AND
        - 2. A medical history which includes previous and current pharmacological and non-pharmacological therapy for the requested diagnosis AND
      - b. The requested agent is NOT prescribed as an as-needed (prn) analgesic AND
      - c. Prescriber has confirmed that a patient-specific pain management plan is on file for the patient AND
      - d. ONE of the following:
        - 1. Patient's medication history includes use of an immediate-acting opioid OR
        - 2. Patient has an intolerance or hypersensitivity to an immediate-acting opioid OR
        - 3. Patient has an FDA labeled contraindication to an immediate-acting opioid AND
      - e. Prescriber has reviewed the patient's records in the state's prescription drug monitoring program (PDMP) AND has determined that the opioid dosages and combinations of opioids and other controlled substances within the patient's records do NOT indicate the patient is at high risk for overdose AND
      - f. Patient does NOT have any FDA labeled contraindications to the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

Opioids ER PA - Tapentadol

### Drug Name(s)

Nucynta Er

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic cancer-related pain OR
  - B. Patient has a diagnosis of pain due to sickle cell disease OR
  - C. Patient is undergoing treatment of chronic non-cancer pain AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
    - iii. ALL of the following:
      - a. Prescriber has completed a formal, consultative evaluation including BOTH of the following:
        - 1. Diagnosis AND
        - 2. A medical history which includes previous and current pharmacological and non-pharmacological therapy for the requested diagnosis AND
      - b. The requested agent is NOT prescribed as an as-needed (prn) analgesic AND
      - c. Prescriber has confirmed that a patient-specific pain management plan is on file for the patient AND
      - d. ONE of the following:
        - 1. Patient's medication history includes use of an immediate-acting opioid OR
        - 2. Patient has an intolerance or hypersensitivity to an immediate-acting opioid OR
        - 3. Patient has an FDA labeled contraindication to an immediate-acting opioid AND
      - e. Prescriber has reviewed the patient's records in the state's prescription drug monitoring program (PDMP) AND has determined that the opioid dosages and combinations of opioids and other controlled substances within the patient's records do NOT indicate the patient is at high risk for overdose AND
      - f. Patient does NOT have any FDA labeled contraindications to the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

Opioids ER PA - Tramadol

#### Drug Name(s)

Conzip

Tramadol Hcl Er

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic cancer-related pain OR
  - B. Patient has a diagnosis of pain due to sickle cell disease OR
  - C. Patient is undergoing treatment of chronic non-cancer pain AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
    - iii. ALL of the following:
      - a. Prescriber has completed a formal, consultative evaluation including BOTH of the following:
        - 1. Diagnosis AND
        - 2. A medical history which includes previous and current pharmacological and non-pharmacological therapy for the requested diagnosis AND
      - b. The requested agent is NOT prescribed as an as-needed (prn) analgesic AND
      - c. Prescriber has confirmed that a patient-specific pain management plan is on file for the patient AND
      - d. ONE of the following:
        - 1. Patient's medication history includes use of an immediate-acting opioid OR
        - 2. Patient has an intolerance or hypersensitivity to an immediate-acting opioid OR
        - 3. Patient has an FDA labeled contraindication to an immediate-acting opioid AND
      - e. Prescriber has reviewed the patient's records in the state's prescription drug monitoring program (PDMP) AND has determined that the opioid dosages and combinations of opioids and other controlled substances within the patient's records do NOT indicate the patient is at high risk for overdose AND
      - f. Patient does NOT have any FDA labeled contraindications to the requested agent

### Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Opzelura PA

### Drug Name(s)

Opzelura

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. ONE of the following:
  - A. Patient (pt) has a diagnosis of mild to moderate atopic dermatitis AND ALL of the following:
    - i. The requested agent will be used for short-term and non-continuous chronic treatment AND
    - ii. Pt is NOT immunocompromised AND
    - iii. ONE of the following:
      - a. Pt has tried and failed a topical steroid (e.g., triamcinolone) OR
      - b. Pt has an intolerance or hypersensitivity to a topical steroid OR
      - c. Pt has an FDA labeled contraindication to a topical steroid AND
    - iv. ONE of the following:
      - a. Pt has tried and failed a topical calcineurin inhibitor (e.g., tacrolimus) OR
      - b. Pt has an intolerance or hypersensitivity to a topical calcineurin inhibitor OR
      - c. Pt has an FDA labeled contraindication to a topical calcineurin inhibitor OR
  - B. Pt has a diagnosis of nonsegmental vitiligo AND
- 2. Pt will NOT be using the requested agent in combination with other biologic immunomodulator agents, other JAK inhibitors, OR potent immunosuppressants (e.g., azathioprine, cyclosporine) AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Pt has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Pt has a diagnosis of mild to moderate atopic dermatitis AND BOTH of the following:
    - i. The requested agent will be used for short-term and non-continuous chronic treatment AND
    - ii. Pt is NOT immunocompromised OR
  - B. Pt has a diagnosis of nonsegmental vitiligo AND
- 3. Pt has had clinical benefit with the requested agent AND
- 4. Pt will NOT be using the requested agent in combination with other biologic immunomodulator agents, other JAK inhibitors, OR potent immunosuppressants (e.g., azathioprine, cyclosporine) AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

### **Age Restriction:**

Patient is 12 years of age or over

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, dermatologist, immunologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis **Coverage Duration:** 

Initial and Renewal: 3 months for atopic dermatitis, other diagnosis - see Other Criteria Other Criteria:

Initial: 6 months for nonsegmental vitiligo, Renewal: 12 months for nonsegmental vitiligo

Oral Immunotherapy Agents PA – Grastek

### Drug Name(s)

Grastek

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of allergic rhinitis, with or without conjunctivitis AND
- 2. Patient's diagnosis is confirmed with ONE of the following:
  - A. Positive skin test to ONE of the pollen extracts included in the requested agent OR
  - B. IgE specific antibodies to ONE of the extracts included in the requested agent: Timothy grass or cross-reactive grass AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to an intranasal corticosteroid AND one other standard allergy agent (e.g., oral or intranasal antihistamines, oral or intranasal corticosteroids, leukotriene inhibitors, note: two separate intranasal corticosteroids meet this criteria) OR
  - B. Patient has an intolerance or hypersensitivity to therapy with an intranasal corticosteroid AND one other standard allergy agent OR
  - C. Patient has an FDA labeled contraindication to therapy with an intranasal corticosteroid AND one other standard allergy agent AND
- 4. Patient will NOT be using the requested agent in combination with a subcutaneous injectable immunotherapy agent AND
- 5. The requested agent will be started, or has already been started, 3 to 4 months before the expected onset of the applicable pollen season AND
- 6. The first dose is given in the clinic/hospital under direct supervision from the provider for a period of at least 30 minutes AND
- 7. Patient has been prescribed epinephrine auto-injector for at home emergency use

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, immunologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be for 12 months

Oral Immunotherapy Agents PA - Odactra

#### Drug Name(s)

Odactra

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of allergic rhinitis, with or without conjunctivitis AND
- 2. Patient's diagnosis is confirmed with ONE of the following:
  - A. Positive skin test to licensed house dust mite allergen extracts OR
  - B. IgE specific antibodies to ONE of the extracts included in the requested agent:
  - Dermatophagoides farinae or Dermatophagoides pteronyssinus AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to an intranasal corticosteroid AND one other standard allergy agent (e.g., oral or intranasal antihistamines, oral or intranasal corticosteroids, leukotriene inhibitors, note: two separate intranasal corticosteroids meet this criteria) OR
  - B. Patient has an intolerance or hypersensitivity to therapy with an intranasal corticosteroid AND one other standard allergy agent OR
  - C. Patient has an FDA labeled contraindication to therapy with an intranasal corticosteroid AND one other standard allergy agent AND
- 4. Patient will NOT be using the requested agent in combination with a subcutaneous injectable immunotherapy agent AND
- 5. The first dose is given in the clinic/hospital under direct supervision from the provider for a period of at least 30 minutes AND
- 6. Patient has been prescribed epinephrine auto-injector for at home emergency use

#### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, immunologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Oral Immunotherapy Agents PA – Oralair

### Drug Name(s)

Oralair

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of allergic rhinitis, with or without conjunctivitis AND
- 2. Patient's diagnosis is confirmed with ONE of the following:
  - A. Positive skin test to ONE of the pollen extracts included in the requested agent OR
  - B. IgE specific antibodies to ONE of the extracts included in the requested agent: Sweet vernal, orchard, perennial rye, Timothy, or Kentucky blue grass AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to an intranasal corticosteroid AND one other standard allergy agent (e.g., oral or intranasal antihistamines, oral or intranasal corticosteroids, leukotriene inhibitors, note: two separate intranasal corticosteroids meet this criteria) OR
  - B. Patient has an intolerance or hypersensitivity to therapy with an intranasal corticosteroid AND one other standard allergy agent OR
  - C. Patient has an FDA labeled contraindication to therapy with an intranasal corticosteroid AND one other standard allergy agent AND
- 4. Patient will NOT be using the requested agent in combination with a subcutaneous injectable immunotherapy agent AND
- 5. The requested agent will be started, or has already been started, 3 to 4 months before the expected onset of the applicable pollen season AND
- 6. The first dose is given in the clinic/hospital under direct supervision from the provider for a period of at least 30 minutes AND
- 7. Patient has been prescribed epinephrine auto-injector for at home emergency use

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, immunologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be for 12 months

Oriahnn PA

### Drug Name(s)

Oriahnn

Indications:

All FDA-Approved Indications.

Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of heavy menstrual bleeding associated with uterine leiomyomas (fibroids) AND
- 2. Patient's diagnosis of uterine fibroids was confirmed via imaging (e.g., ultrasound) AND
- 3. Patient is premenopausal AND
- 4. Patient will NOT be using the requested agent in combination with another GnRH antagonist agent [e.g., Orilissa (elagolix), Myfembree (relugolix, estradiol hemihydrate, norethindrone acetate)] for the requested indication AND
- 5. The requested dose is within FDA labeled dosing for the requested indication AND
- 6. ONE of the following:
  - A. Patient is initiating therapy with the requested agent OR
  - B. BOTH of the following:
    - i. Patient is continuing therapy with the requested agent and the prescriber has provided information indicating the number of months the patient has been on therapy with the requested agent AND
    - ii. The total duration of treatment with the requested agent has NOT exceeded 24 months per lifetime

### Age Restriction:

### **Prescriber Restrictions:**

## **Coverage Duration:**

For no prior Oriahnn use: 24 months

For Prior Oriahnn use: Remainder of 24 months

Orilissa PA

### Drug Name(s)

Orilissa

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of moderate to severe pain associated with endometriosis AND
- 2. ONE of the following:
  - A. Patient has coexisting dyspareunia AND has not received 6 or more months of therapy with the requested agent OR
  - B. Patient has coexisting moderate hepatic impairment (Child-Pugh Class B) AND has not received 6 or more months of therapy with the requested agent OR
  - C. Patient does not have coexisting dyspareunia or moderate hepatic impairment (Child-Pugh Class B), AND has not received 24 or more months of therapy with the requested agent AND
- 3. Patient will NOT be using the requested agent in combination with another GnRH antagonist agent [e.g., Oriahnn (elagolix, estradiol, norethindrone acetate), Myfembree (relugolix, estradiol hemihydrate, norethindrone acetate)] for the requested indication AND
- 4. The requested dose is within FDA labeled dosing for the requested indication AND
- 5. ONE of the following:
  - A. Patient is initiating therapy with the requested agent OR
  - B. BOTH of the following:
    - i. Patient is continuing therapy with the requested agent and the prescriber has provided information indicating the number of months the patient has been on therapy with the requested agent AND
    - ii. The total duration of treatment with the requested agent has NOT exceeded 6 months per lifetime if patient has coexisting dyspareunia or moderate hepatic impairment, or 24 months per lifetime if the patient does NOT have coexisting dyspareunia or moderate hepatic impairment

#### Age Restriction:

#### **Prescriber Restrictions:**

## **Coverage Duration:**

For no prior Orilissa use and for prior Orilissa use, see Other Criteria for approval

#### Other Criteria:

No prior Orilissa use: Approve 6 months for coexisting dyspareunia OR moderate hepatic impairment, Approve 24 months for no coexisting condition

Prior Orilissa use: Approve remainder of 6 months for coexisting dyspareunia OR moderate hepatic impairment, Approve remainder of 24 months for no coexisting condition

Orkambi PA

### Drug Name(s)

Orkambi

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of cystic fibrosis AND
- 2. ONE of the following:
  - A. Patient has the presence of the F508del mutation on both alleles (homozygous) of the CFTR gene confirmed by genetic testing OR
  - B. Patient has another CFTR gene mutation(s) that is responsive to the requested agent, as indicated in the FDA label, confirmed by genetic testing AND
- 3. Patient will NOT be using the requested agent in combination with another CFTR modulator agent for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of cystic fibrosis AND
- 3. Patient has had improvement or stabilization with the requested agent [e.g., improvement in FEV1 from baseline, increase in weight/BMI, improvement from baseline Cystic Fibrosis Questionnaire-Revised (CFQ-R) Respiratory Domain score, improvements in respiratory symptoms (cough, sputum production, and difficulty breathing), and/or reduced number of pulmonary exacerbations] AND
- 4. Patient will NOT be using the requested agent in combination with another CFTR modulator agent for the requested indication

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cystic fibrosis, pulmonologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be for 12 months

Osphena PA

Drug Name(s)

Osphena

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Otezla PA

#### Drug Name(s)

Otezla

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. Patient has ONE of the following diagnoses:
      - 1. Plaque psoriasis OR
      - 2. Active psoriatic arthritis AND
    - ii. ONE of the following:
      - 1. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - 2. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
      - 3. Patient's medication history indicates use of a biologic immunomodulator agent for the same FDA labeled indication OR
      - 4. Patient has tried and had an inadequate response to at least ONE conventional prerequisite agent for the requested indication OR
      - 5. Patient has an intolerance or hypersensitivity to at least ONE conventional prerequisite agent for the requested indication OR
      - 6. Patient has an FDA labeled contraindication to at least ONE conventional prerequisite agent for the requested indication OR
  - B. Patient has a diagnosis of oral ulcers associated with Behcet's disease (BD) AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has ONE of the following diagnoses:
  - A. Plaque psoriasis OR
  - B. Active psoriatic arthritis OR
  - C. Oral ulcers associated with Behcet's disease (BD) AND
- 3. Patient has had clinical benefit with the requested agent (slowing of disease progression or decrease in symptom severity and/or frequency) AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

#### **Prescriber Restrictions:**

## **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Formulary conventional agent required for diagnoses of plaque psoriasis or active psoriatic arthritis

Formulary conventional agents for plaque psoriasis include cyclosporine, methotrexate, tazarotene, topical calcitriol, or topical corticosteroids

Formulary conventional agents for active psoriatic arthritis include cyclosporine, leflunomide, methotrexate, or sulfasalazine

NO prerequisites are required for a diagnosis of oral ulcers associated with Behcet's disease (BD)

Otrexup PA

## Drug Name(s)

Otrexup

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. Patient has tried and had an inadequate response to a generic methotrexate injectable agent OR
  - B. Patient has an intolerance or hypersensitivity to a generic methotrexate injectable agent OR
  - C. Patient has an FDA labeled contraindication to a generic methotrexate injectable agent OR
  - D. Prescriber has provided information that the patient has a physical or a mental disability that would prevent the patient from using a generic methotrexate injectable agent

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Oxbryta PA

## Drug Name(s)

Oxbryta

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of sickle cell disease AND
- 2. ONE of the following:
  - A. Patient has tried and had an inadequate response to maximally tolerated dose of hydroxyurea OR
  - B. Patient has an intolerance or hypersensitivity to hydroxyurea OR
  - C. Patient has an FDA labeled contraindication to hydroxyurea

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of sickle cell disease AND
- 3. Patient has had clinical benefit with the requested agent

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Oxervate PA

## Drug Name(s)

Oxervate

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of neurotrophic keratitis (NK) AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., ophthalmologist, optometrist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 8 weeks

Oxlumo PA

### Drug Name(s)

Oxlumo

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of primary hyperoxaluria type 1 (PH1) confirmed by ONE of the following:
  - A. Genetic testing of the AGXT gene indicates a pathogenic mutation OR
  - B. Liver biopsy demonstrates absent or significantly reduced alanine:glyoxylate aminotransferase (AGT) activity AND
- 2. The requested agent will be used to lower urinary or plasma oxalate levels AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of primary hyperoxaluria type 1 (PH1) AND
- 3. The requested agent will be used to lower urinary or plasma oxalate levels AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, nephrologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be 6 months for initial, 12 months for renewal

Ozempic PA

### Drug Name(s)

Ozempic

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

Requested agent will be used for weight loss alone

### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of type 2 diabetes mellitus AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 180 days OR
  - C. ALL of the following:
    - i. ONE of the following:
      - 1. Patient's medication history includes use of a non glucagon-like peptide-1 (GLP-1) oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) within the past 90 days OR
      - 2. Patient had an ineffective treatment response to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 3. Patient has an intolerance or hypersensitivity to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 4. Patient has an FDA labeled contraindication to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 5. BOTH of the following:
        - a. Patient has a diagnosis of established cardiovascular disease [e.g., myocardial infarction, stroke, any revascularization procedure, transient ischemic attack, unstable angina, amputation, symptomatic or asymptomatic coronary artery disease] AND
        - b. The requested agent will be used to reduce the risk of major adverse cardiovascular events (cardiovascular death, non-fatal myocardial infarction, or non-fatal stroke) AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent AND
    - iii. Patient will NOT be using the requested agent in combination with another GLP-1 agonist agent, or an agent containing a GLP-1 agonist AND
    - iv. Patient will NOT be using the requested agent in combination with an agent containing a dipeptidyl peptidase-4 (DPP-4) inhibitor

**Age Restriction:** 

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

Palforzia PA

### Drug Name(s)

Palforzia

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of peanut allergy AND
- 2. Patient is/was 4-17 years of age at the time of initiating therapy AND
- 3. Patient has been prescribed epinephrine injection for at home emergency use AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, immunologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Palynziq PA

## Drug Name(s)

Palynziq

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of phenylketonuria (PKU) AND
- 2. Patient has a baseline blood Phe level greater than 600 micromol/L (10 mg/dL) AND
- 3. Patient will NOT be using the requested agent in combination with sapropterin for the requested indication AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of phenylketonuria (PKU) AND
- 3. ONE of the following:
  - a. Patient's blood Phe levels are being maintained within the acceptable range OR
  - b. Patient has had a decrease in blood Phe level from baseline AND
- 4. Patient will NOT be using the requested agent in combination with sapropterin for the requested indication AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., metabolic or genetic disorders) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be 9 months for initial, 12 months for renewal

Panretin PA

### Drug Name(s)

Panretin

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of cutaneous lesions associated with AIDS-related Kaposi's sarcoma (KS) OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. ALL of the following:
    - i. ONE of the following:
      - 1. BOTH of the following:
        - a. Patient has a diagnosis of cutaneous lesions associated with AIDS-related Kaposi's sarcoma (KS) AND
        - b. Patient does NOT require systemic anti-Kaposi's sarcoma therapy OR
      - 2. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Prescriber is a specialist in the area of the patient's diagnosis (e.g., oncologist, dermatologist, infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis AND
    - iii. Patient does NOT have any FDA labeled contraindications to the requested agent

## Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Panzyga PA

Drug Name(s)

Panzyga

Indications:

All Medically-Accepted Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Pegylated Interferon PA

### Drug Name(s)

Pegasys

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chronic hepatitis B AND BOTH of the following:
    - i. The chronic hepatitis B infection has been confirmed by serological markers AND
    - ii. Patient has NOT been administered the requested agent for more than 48 weeks for the treatment of chronic hepatitis B OR
  - B. BOTH of the following:
    - i. Patient has a diagnosis of chronic hepatitis C confirmed by serological markers AND
    - ii. The requested agent will be used in a treatment regimen and length of therapy that is supported in FDA approved labeling for the patient's diagnosis and genotype OR
  - C. Patient has an indication that is supported in CMS approved compendia for the requested agent

### Age Restriction:

#### **Prescriber Restrictions:**

### **Coverage Duration:**

12 months for all other diagnoses. For hep B, hep C see Other Criteria

#### Other Criteria:

No prior peginterferon alfa use, approve 48 weeks for hepatitis B infection. Prior peginterferon alfa use, approve remainder of 48 weeks of total therapy for hepatitis B infection

Duration of therapy for hepatitis C: Based on FDA approved labeling

Pirfenidone PA

## Drug Name(s)

**Esbriet** 

Pirfenidone

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of idiopathic pulmonary fibrosis (IPF) AND
- 2. Patient has no known explanation for interstitial lung disease (ILD) or pulmonary fibrosis (e.g., radiation, drugs, metal dusts, sarcoidosis, or any connective tissue disease known to cause ILD)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of idiopathic pulmonary fibrosis (IPF) AND
- 3. Patient has had clinical benefit with the requested agent

### Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., pathologist, pulmonologist, radiologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be for 12 months

Posaconazole PA

### Drug Name(s)

Noxafil

Posaconazole Inj

Posaconazole Dr

Posaconazole Susp

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

#### **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of oropharyngeal candidiasis AND ONE of the following:
    - i. Patient has tried and had an inadequate response to fluconazole or an alternative antifungal agent OR
    - ii. Patient has an intolerance or hypersensitivity to fluconazole or an alternative antifungal agent OR
    - iii. Patient has an FDA labeled contraindication to fluconazole or an alternative antifungal agent OR
  - B. The requested agent is being prescribed for prophylaxis of invasive Aspergillus or Candida AND patient is severely immunocompromised, such as a hematopoietic stem cell transplant [HSCT] recipient, or hematologic malignancy with prolonged neutropenia from chemotherapy, or is a high-risk solid organ (lung, heart-lung, liver, pancreas, small bowel) transplant patient, or long term use of high dose corticosteroids (greater than 1 mg/kg/day of prednisone or equivalent) OR
  - C. Patient has a diagnosis of invasive Aspergillus AND ONE of the following:
    - i. Patient has tried and had an inadequate response to an alternative antifungal agent OR
    - ii. Patient has an intolerance or hypersensitivity to an alternative antifungal agent OR
    - iii. Patient has an FDA labeled contraindication to an alternative antifungal agent OR
  - D. Patient has another indication that is supported in CMS approved compendia for the requested agent

### Age Restriction:

#### **Prescriber Restrictions:**

### **Coverage Duration:**

One month for oropharyngeal candidiasis, 6 months for all other indications

#### Other Criteria:

Criteria for renewal approval require BOTH of the following:

1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND

### 2. ONE of the following:

- A. The requested agent is being prescribed for prophylaxis of invasive Aspergillus or Candida and patient continues to be severely immunocompromised, such as a hematopoietic stem cell transplant [HSCT] recipient, or hematologic malignancy with prolonged neutropenia from chemotherapy, or is a high-risk solid organ (lung, heart-lung, liver, pancreas, small bowel) transplant patient, or long term use of high dose corticosteroids (greater than 1 mg/kg/day of prednisone or equivalent) OR
- B. Patient has a diagnosis of invasive Aspergillus AND patient has continued indicators of active disease (e.g., continued radiologic findings, positive cultures, positive serum galactomannan assay for Aspergillus) OR
- C. BOTH of the following:
  - i. Patient has a diagnosis of oropharyngeal candidiasis AND
  - ii. Patient has had clinical benefit with the requested agent OR
- D. BOTH of the following:
  - i. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
  - ii. Patient has had clinical benefit with the requested agent

Praluent PA

Drug Name(s)

Praluent

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Prolia PA

### Drug Name(s)

Prolia

#### Indications:

All FDA-Approved Indications, Some Medically-Accepted Indications.

#### Off-Label Uses:

Osteopenia (osteoporosis prophylaxis)

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of:

#### 1. ONE of:

A. Patient's (pt) sex is male or the pt is postmenopausal with a diagnosis of osteoporosis AND BOTH of:

- i. Pt's diagnosis was confirmed by ONE of:
  - 1. A fragility fracture in the hip or spine OR
  - 2. A T-score of -2.5 or lower OR
  - 3. A T-score of -1.0 to -2.5 AND ONE of:
    - a. A fragility fracture of the proximal humerus, pelvis, or distal forearm OR
    - b. A FRAX 10-year probability for major osteoporotic fracture of 20% or greater  $\mathsf{OR}$
    - c. A FRAX 10-year probability of hip fracture of 3% or greater AND

#### ii. ONE of:

- 1. Pt is at a very high fracture risk as defined by ONE of:
  - a. Pt had a recent fracture (within the past 12 months) OR
  - b. Pt had fractures while on FDA approved osteoporosis therapy OR
  - c. Pt has had multiple fractures OR
  - d. Pt had fractures while on drugs causing skeletal harm (e.g., long-term glucocorticoids) OR
  - e. Pt has a very low T-score (less than -3.0) OR
  - f. Pt is at high risk for falls or has a history of injurious falls OR
  - g. Pt has a very high fracture probability by FRAX (e.g., major osteoporosis fracture greater than 30%, hip fracture greater than 4.5%) or by other validated fracture risk algorithm OR
- 2. ONE of:
  - a. Pt's medication history includes use of a bisphosphonate OR
  - b. Pt has an intolerance, FDA labeled contraindication, or hypersensitivity to a bisphosphonate OR
- B. Pt is requesting the agent for osteopenia (osteoporosis prophylaxis) AND ALL of:
  - i. ONE of:
    - 1. Pt's sex is male and the pt is 50 years of age or over OR
    - 2. Pt is postmenopausal AND

- ii. Pt has a T-score between -1.0 to -2.50 AND
- iii. ONE of:
  - a. A fragility fracture of the proximal humerus, pelvis, or distal forearm OR
  - b. 10-year probability of a hip fracture 3% and greater per FRAX OR
  - c. 10-year probability of a major OP-related fracture 20% and greater per FRAX AND
- iv. ONE of:
  - a. Pt's medication history includes use of a bisphosphonate OR

Criteria continues: See Other Criteria

Age Restriction:

Prescriber Restrictions: Coverage Duration:

Approval will be for 12 months

- b. Pt has an intolerance, FDA labeled contraindication, or hypersensitivity to a bisphosphonate OR
- C. Pt's sex is a female with a diagnosis of breast cancer who is receiving aromatase inhibitor therapy AND ONE of:
  - i. Pt's medication history includes use of a bisphosphonate OR
  - ii. Pt has an intolerance, FDA labeled contraindication, or hypersensitivity to a bisphosphonate OR
- D. Pt's sex is male with a diagnosis of prostate cancer receiving androgen deprivation therapy (ADT) AND ONE of:
  - i. Pt's medication history includes use of a bisphosphonate OR
  - ii. Pt has an intolerance, FDA labeled contraindication, or hypersensitivity to a bisphosphonate OR
- E. Pt has a diagnosis of glucocorticoid-induced osteoporosis AND ALL of:
  - i. Pt is either initiating or continuing systemic glucocorticoids in a daily dose equivalent to 7.5 mg or greater of prednisone AND
  - ii. Pt is expected to remain on glucocorticoids for at least 6 months AND
  - iii. Pt's diagnosis was confirmed by ONE of:
    - 1. A fragility fracture in the hip or spine OR
    - 2. A T-score of -2.5 or lower OR
    - 3. A T-score of -1.0 to -2.5 AND ONE of the following:
      - a. A fragility fracture of the proximal humerus, pelvis, or distal forearm
      - b. A FRAX 10-year probability for major osteoporotic fracture of 20% or greater OR
      - c. A FRAX 10-year probability of hip fracture of 3% or greater AND
  - iv. ONE of:
    - 1. Pt is at a very high fracture risk as defined by ONE of the following:
      - a. Pt had a recent fracture (within the past 12 months) OR
      - b. Pt had fractures while on FDA approved osteoporosis therapy OR

- c. Pt has had multiple fractures OR
- d. Pt had fractures while on drugs causing skeletal harm (e.g., long-term glucocorticoids) OR
- e. Pt has a very low T-score (less than -3.0) OR
- f. Pt is at high risk for falls or has a history of injurious falls OR
- g. Pt has a very high fracture probability by FRAX (e.g., major osteoporosis fracture greater than 30%, hip fracture greater than 4.5%) or by other validated fracture risk algorithm OR

#### 2. ONE of:

- a. Pt's medication history includes use of a bisphosphonate OR
- b. Pt has an intolerance, FDA labeled contraindication, or hypersensitivity to a bisphosphonate AND

#### 2. ONE of:

- A. Pt has a pretreatment or current calcium level that is NOT below the lower limit of the testing laboratory's normal range OR
- B. Pt has a pretreatment or current calcium level that is below the lower limit of the testing laboratory's normal range AND it will be corrected prior to use of the requested agent OR
- C. Prescriber has indicated that the pt is not at risk for hypocalcemia (not including risk associated with the requested agent) AND
- 3. Pt will NOT be using the requested agent in combination with a bisphosphonate, another form of denosumab (e.g., Xgeva), romosozumab-aqqg, or parathyroid hormone analog (e.g., abaloparatide, teriparatide) for the requested indication AND
- 4. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Promacta PA

## Drug Name(s)

Promacta

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient (pt) has a diagnosis of persistent or chronic immune (idiopathic) thrombocytopenia (ITP) AND ONE of the following:
    - i. Pt has tried and had an insufficient response to a corticosteroid or immunoglobulin (IVIg or anti-D) OR
    - ii. Pt has an intolerance or hypersensitivity to a corticosteroid or immunoglobulin (IVIg or anti-D) OR
    - iii. Pt has an FDA labeled contraindication to a corticosteroid or immunoglobulin (IVIg or anti-D) OR
    - iv. Pt has had an insufficient response to a splenectomy OR
  - B. Pt has a diagnosis of hepatitis C associated thrombocytopenia AND ONE of the following:
    - i. Pt's platelet count is less than 75 x  $10^9$ /L AND the intent is to increase platelet counts sufficiently to initiate interferon therapy OR
    - ii. Pt is on concomitant therapy with interferon therapy AND is at risk for discontinuing hepatitis C therapy due to thrombocytopenia OR
  - C. Pt has a diagnosis of severe aplastic anemia (SAA) AND ALL of the following:
    - i. Pt has at least 2 of the following blood criteria:
      - 1. Neutrophils less than 0.5 X 10<sup>9</sup>/L OR
      - 2. Platelets less than 30 X 10<sup>9</sup>/L OR
      - 3. Reticulocyte count less than 60 X 10^9/L AND
    - ii. Pt has at least 1 of the following marrow criteria:
      - 1. Severe hypocellularity is less than 25% OR
      - 2. Moderate hypocellularity is 25-50% with hematopoietic cells representing less than 30% of residual cells AND
    - iii. ONE of the following:
      - 1. Pt has tried and had an insufficient response to BOTH antithymocyte globulin (ATG) AND cyclosporine therapy OR
      - 2. BOTH of the following:
        - a. Pt will be using the requested agent as first-line treatment (i.e., has not been treated with ATG and/or cyclosporine) AND
        - b. Pt will use the requested agent in combination with standard immunosuppressive therapy (i.e., ATG AND cyclosporine) OR

Initial criteria continues: see Other Criteria

Age Restriction:

Prescriber Restrictions: Coverage Duration:

Initial: 6 months for ITP. Renewal: 12 months for ITP. Other indications, see Other Criteria.

#### Other Criteria:

- D. Pt has another indication that is supported in CMS approved compendia for the requested agent AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Pt has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Pt has a diagnosis of persistent or chronic immune (idiopathic) thrombocytopenia (ITP) AND ONE of the following:
    - i. Pt's platelet count is 50 x 10^9/L or greater OR
    - ii. Pt's platelet count has increased sufficiently to avoid clinically significant bleeding OR
  - B. Pt has a diagnosis of hepatitis C associated thrombocytopenia AND BOTH of the following:
    - i. ONE of the following:
      - 1. Pt will be initiating hepatitis C therapy with interferon therapy OR
      - 2. Pt will be maintaining hepatitis C therapy with interferon therapy at the same time as the requested agent AND
    - ii. ONE of the following:
      - 1. Pt's platelet count is 90 x 10^9/L or greater OR
      - 2. Pt's platelet count has increased sufficiently to initiate or maintain interferon therapy for the treatment of hepatitis C OR
  - C. Pt has a diagnosis of severe aplastic anemia (SAA) AND the pt has had clinical benefit with the requested agent OR
  - D. Pt has another indication that is supported in CMS approved compendia and the pt has had clinical benefit with the requested agent AND
- 3. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Initial: 48 weeks for hepatitis C associated thrombocytopenia, 6 months for first-line therapy in severe aplastic anemia, 16 weeks for SAA, 12 months for All other indications

Renewal: 48 weeks for hepatitis C associated thrombocytopenia, 12 months for SAA, 12 months for All other indications

Pulmonary Hypertension PA – Adempas

## Drug Name(s)

Adempas

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent within the past 90 days AND
    - ii. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has a diagnosis of chronic thromboembolic pulmonary hypertension (CTEPH), WHO Group 4, as determined by a ventilation-perfusion scan and a confirmatory selective pulmonary angiography AND ALL of the following:
    - i. ONE of the following:
      - a. Patient is NOT a candidate for surgery OR
      - b. Patient has had pulmonary endarterectomy AND has persistent or recurrent disease AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
  - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units OR C. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1 as determined by right heart catheterization AND ALL of the following:
    - i. Patient's World Health Organization (WHO) functional class is II or greater AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
    - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units AND

Initial criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

#### Other Criteria:

- v. ONE of the following:
  - a. The requested agent will be utilized as monotherapy OR
  - b. The requested agent will be utilized for add-on therapy to existing monotherapy (dual therapy), AND BOTH of the following:
    - 1. Patient has unacceptable or deteriorating clinical status despite established pharmacotherapy AND
    - 2. The requested agent is in a different therapeutic class OR
  - c. The requested agent will be utilized for add-on therapy to existing dual therapy (triple therapy), AND ALL of the following:
    - 1. ONE of the following:
      - i. A prostanoid has been started as one of the agents in the triple therapy OR
      - ii. Patient has an intolerance or hypersensitivity to a prostanoid OR
      - iii. Patient has an FDA labeled contraindication to a prostanoid AND
    - 2. Patient has unacceptable or deteriorating clinical status despite established pharmacotherapy AND
    - 3. All three agents in the triple therapy are from a different therapeutic class

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Pulmonary Hypertension PA – Ambrisentan

## Drug Name(s)

Ambrisentan

Letairis

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent within the past 90 days AND
    - ii. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1 as determined by right heart catheterization AND ALL of the following:
    - i. Patient's World Health Organization (WHO) functional class is II or greater AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
    - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units
    - v. ONE of the following:
      - a. The requested agent will be utilized as monotherapy OR
      - b. The requested agent will be used in combination with a phosphodiesterase 5 (PDE5) inhibitor for dual therapy ONLY OR
      - c. The requested agent will be utilized for add-on therapy to existing monotherapy (dual therapy), [except for dual therapy requests for a phosphodiesterase 5 (PDE 5) inhibitor plus an endothelin receptor antagonist (ERA)], AND BOTH of the following:
        - 1. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 2. The requested agent is in a different therapeutic class OR

Initial criteria continues: see Other Criteria

Age Restriction:

Prescriber Restrictions: Coverage Duration:

#### Approval will be for 12 months

## Other Criteria:

- d. The requested agent will be utilized for add-on therapy to existing dual therapy (triple therapy), AND ALL of the following:
  - 1. ONE of the following:
    - i. A prostanoid has been started as one of the agents in the triple therapy OR
    - ii. Patient has an intolerance or hypersensitivity to a prostanoid OR
    - iii. Patient has an FDA labeled contraindication to a prostanoid
  - 2. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
  - 3. All three agents in the triple therapy are from a different therapeutic class OR
- e. The requested agent will be utilized as part of triple therapy in a treatment naive patient AND BOTH of the following:
  - 1. Patient is classified as WHO functional class IV AND
  - 2. The three agents being utilized consist of: ERA plus PDE5i plus prostanoid

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Pulmonary Hypertension PA – Bosentan

#### Drug Name(s)

Bosentan

Tracleer

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

Elevated liver enzymes accompanied by signs or symptoms of liver dysfunction/injury or a bilirubin level of 2 times the ULN (upper limit of normal) or greater AND FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent within the past 90 days AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1, as determined by right heart catheterization, AND ALL of the following:
    - i. Patient's World Health Organization (WHO) functional class is II or greater AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
    - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units AND
    - v. ONE of the following:
      - a. The requested agent will be utilized as monotherapy OR
      - b. The requested agent will be used in combination with a phosphodiesterase 5 (PDE5) inhibitor for dual therapy ONLY OR
      - c. The requested agent will be utilized for add-on therapy to existing monotherapy (dual therapy), [except for dual therapy requests for a phosphodiesterase 5 inhibitor (PDE5) plus an endothelin receptor antagonist (ERA)], AND BOTH of the following:
        - 1. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 2. The requested agent is in a different therapeutic class OR

Initial criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

- d. The requested agent will be utilized for add-on therapy to existing dual therapy (triple therapy), AND ALL of the following:
  - 1. ONE of the following:
    - i. A prostanoid has been started as one of the agents in the triple therapy OR
    - ii. Patient has an intolerance or hypersensitivity to a prostanoid OR
    - iii. Patient has an FDA labeled contraindication to a prostanoid AND
  - 2. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
  - 3. All three agents in the triple therapy are from a different therapeutic class OR
- e. The requested agent will be utilized as part of triple therapy in a treatment naive patient AND BOTH of the following:
  - 1. Patient is classified as WHO functional class IV AND
  - 2. The three agents being utilized consist of: ERA plus PDE5i plus prostanoid OR
- C. Patient has an indication that is supported in CMS approved compendia for the requested agent

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Pulmonary Hypertension PA – Opsumit

## Drug Name(s)

Opsumit

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent within the past 90 days AND
    - ii. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1 as determined by right heart catheterization AND ALL of the following:
    - i. Patient's World Health Organization (WHO) functional class is II or greater AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
    - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units AND
    - v. ONE of the following:
      - a. The requested agent will be utilized as monotherapy OR
      - b. The requested agent will be used in combination with a phosphodiesterase 5 (PDE5) inhibitor for dual therapy ONLY OR
      - c. The requested agent will be utilized for add-on therapy to existing monotherapy (dual therapy), [except for dual therapy requests for a phosphodiesterase 5 (PDE 5) inhibitor plus an endothelin receptor antagonist (ERA)], AND BOTH of the following:
        - 1. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 2. The requested agent is in a different therapeutic class OR

Initial criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

#### Other Criteria:

- d. The requested agent will be utilized for add-on therapy to existing dual therapy (triple therapy), AND ALL of the following:
  - 1. ONE of the following:
    - i. A prostanoid has been started as one of the agents in the triple therapy OR
    - ii. Patient has an intolerance or hypersensitivity to a prostanoid OR
    - iii. Patient has an FDA labeled contraindication to a prostanoid AND
  - 2. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
  - 3. All three agents in the triple therapy are from a different therapeutic class OR
- e. The requested agent will be utilized as part of triple therapy in a treatment naive patient AND BOTH of the following:
  - 1. Patient is classified as WHO functional class IV AND
  - 2. The three agents being utilized consist of: ERA plus PDE5i plus prostanoid

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Pulmonary Hypertension PA – Opsynvi

# Drug Name(s)

Opsynvi

## **Indications:**

All FDA-Approved Indications.

## Off-Label Uses:

## **Exclusion Criteria:**

Concurrently taking another phosphodiesterase 5 (PDE 5) inhibitor with the requested agent AND FDA labeled contraindications to the requested agent

# **Required Medical Information:**

**Pending CMS Review** 

Pulmonary Hypertension PA – Orenitram

## Drug Name(s)

Orenitram

Orenitram Titration Kit Month 1

Orenitram Titration Kit Month 2

Orenitram Titration Kit Month 3

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

#### **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent within the past 90 days AND
    - ii. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1 as determined by right heart catheterization AND ALL of the following:
    - i. Patient's World Health Organization (WHO) functional class is II or greater AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
    - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units AND
    - v. ONE of the following:
      - a. The requested agent will be utilized as monotherapy OR
      - b. The requested agent will be utilized for add-on therapy to existing monotherapy (dual therapy), AND BOTH of the following:
        - 1. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 2. The requested agent is in a different therapeutic class OR
      - c. The requested agent will be utilized for add-on therapy to existing dual therapy (triple therapy), AND ALL of the following:
        - 1. Patient is WHO functional class III or IV AND
        - 2. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND

- 3. All three agents in the triple therapy are from a different therapeutic class OR
- d. The requested agent will be utilized as part of triple therapy in a treatment naive patient AND BOTH of the following:
  - 1. Patient is classified as WHO functional class IV AND
  - 2. The three agents being utilized consist of: ERA plus PDE5i plus prostanoid

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Pulmonary Hypertension PA - Sildenafil

#### Drug Name(s)

Ligrev

Revatio

Sildenafil Citrate

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

Concurrently taking another phosphodiesterase 5 (PDE 5) inhibitor with the requested agent AND FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent within the past 90 days AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1 as determined by right heart catheterization AND ALL of the following:
    - i. Patient's World Health Organization (WHO) functional class is II or greater AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
    - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units AND
    - v. ONE of the following:
      - a. The requested agent will be utilized as monotherapy OR
      - b. The requested agent will be used in combination with an endothelin receptor antagonist (ERA) for dual therapy ONLY OR
      - c. The requested agent will be utilized for add-on therapy to existing monotherapy, [except for dual requests for a phosphodiesterase 5 (PDE5) inhibitor plus an endothelin receptor antagonist (ERA)], AND BOTH of the following:
        - 1. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 2. The requested agent is in a different therapeutic class OR

Initial criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

- d. The requested agent will be utilized for add-on therapy to existing dual therapy (triple therapy) AND ALL of the following:
  - 1. ONE of the following:
    - i. A prostanoid has been started as one of the agents in the triple therapy OR
    - ii. Patient has an intolerance or hypersensitivity to a prostanoid OR
    - iii. Patient has an FDA labeled contraindication to a prostanoid
  - 2. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
  - 3. All three agents in the triple therapy are from a different therapeutic class OR
- e. The requested agent will be utilized as part of triple therapy in a treatment naive patient AND BOTH of the following:
  - 1. Patient is classified as WHO functional class IV AND
  - 2. The three agents being utilized consist of: ERA plus PDE5i plus prostanoid OR
- C. Patient has an indication that is supported in CMS approved compendia for the requested agent

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Pulmonary Hypertension PA - Tadalafil

## Drug Name(s)

Adcirca

Alva

Tadalafil Tablet 20Mg

Tadliq

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

Concurrently taking another phosphodiesterase 5 (PDE 5) inhibitor with the requested agent AND FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent within the past 90 days AND
    - ii. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent OR
  - B. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1 as determined by right heart catheterization AND ALL of the following:
    - i. Patient's World Health Organization (WHO) functional class is II or greater AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
    - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units AND
    - v. ONE of the following:
      - a. The requested agent will be utilized as monotherapy OR
      - b. The requested agent will be used in combination with an endothelin receptor antagonist (ERA) for dual therapy ONLY OR
      - c. The requested agent will be utilized for add-on therapy to existing monotherapy, [except for dual requests for a phosphodiesterase 5 (PDE5) inhibitor plus an endothelin receptor antagonist (ERA)], AND BOTH of the following:
        - 1. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND

453

2. The requested agent is in a different therapeutic class OR

Initial criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

- d. The requested agent will be utilized for add-on therapy to existing dual therapy (triple therapy) AND ALL of the following:
  - 1. ONE of the following:
    - i. A prostanoid has been started as one of the agents in the triple therapy OR
    - ii. Patient has an intolerance or hypersensitivity to a prostanoid OR
    - iii. Patient has an FDA labeled contraindication to a prostanoid AND
  - 2. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
  - 3. All three agents in the triple therapy are from a different therapeutic class OR
- e. The requested agent will be utilized as part of triple therapy in a treatment naive patient AND BOTH of the following:
  - 1. Patient is classified as WHO functional class IV AND
  - 2. The three agents being utilized consist of: ERA plus PDE5i plus prostanoid OR
- C. Patient has an indication that is supported in CMS approved compendia for the requested agent

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Pulmonary Hypertension PA – Tyvaso DPI

### Drug Name(s)

Tyvaso DPI Institutional Kit

Tyvaso DPI Maintenance Kit

Tyvaso DPI Titration Kit

### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent within the past 90 days AND
    - ii. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1 as determined by right heart catheterization AND ALL of the following:
    - i. Patient's World Health Organization (WHO) functional class is II or greater AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
    - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units AND
    - v. ONE of the following:
      - a. The requested agent will be utilized as monotherapy OR
      - b. The requested agent will be utilized for add-on therapy to existing monotherapy (dual therapy), AND BOTH of the following:
        - 1. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 2. The requested agent is in a different therapeutic class OR
      - c. The requested agent will be utilized for add-on therapy to existing dual therapy (triple therapy), AND ALL of the following:
        - 1. Patient is WHO functional class III or IV AND
        - 2. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 3. All three agents in the triple therapy are from a different therapeutic class OR

- d. The requested agent will be utilized as part of triple therapy in a treatment naive patient AND BOTH of the following:
  - 1. Patient is classified as WHO functional class IV AND
  - 2. The three agents being utilized consist of: ERA plus PDE5i plus prostanoid OR

Initial criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

#### Other Criteria:

C. Patient has a diagnosis of pulmonary hypertension associated with interstitial lung disease (PH-ILD, WHO group 3) as determined by right heart catheterization AND ALL of the following:

- i. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
- ii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
- iii. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units AND
- iv. Patient has a forced vital capacity (FVC) less than 70% of predicted AND
- v. Patient will continue standard of care therapy for ILD (e.g., nintedanib)

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. ONE of the following:
  - A. Patient has a diagnosis of pulmonary hypertension associated with interstitial lung disease (PH-ILD, WHO group 3) AND the patient will continue standard of care therapy for ILD (e.g., nintedanib) OR
  - B. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1

Pulmonary Hypertension PA – Uptravi

## Drug Name(s)

Uptravi

Uptravi Titration Pack

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent within the past 90 days AND
    - ii. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1 as determined by right heart catheterization AND ALL of the following:
    - i. Patient's World Health Organization (WHO) functional class is II or greater AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
    - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units
    - v. ONE of the following:
      - a. The requested agent will be utilized as monotherapy OR
      - b. The requested agent will be utilized for add-on therapy to existing monotherapy (dual therapy), AND BOTH of the following:
        - 1. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 2. The requested agent is in a different therapeutic class OR
      - c. The requested agent will be utilized for add-on therapy to existing dual therapy (triple therapy), AND BOTH of the following:
        - 1. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 2. All three agents in the triple therapy are from a different therapeutic

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Pulmonary Hypertension PA – Ventavis

## Drug Name(s)

Ventavis

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. ONE of the following:
      - a. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
      - b. Prescriber states the patient is currently being treated with the requested agent within the past 90 days AND
    - ii. Patient has an FDA labeled indication for the requested agent OR
  - B. Patient has a diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1 as determined by right heart catheterization AND ALL of the following:
    - i. Patient's World Health Organization (WHO) functional class is II or greater AND
    - ii. Patient has a mean pulmonary arterial pressure greater than 20 mmHg AND
    - iii. Patient has a pulmonary capillary wedge pressure less than or equal to 15 mmHg AND
    - iv. Patient has a pulmonary vascular resistance greater than or equal to 3 Wood units AND
    - v. ONE of the following:
      - a. The requested agent will be utilized as monotherapy OR
      - b. The requested agent will be utilized for add-on therapy to existing monotherapy (dual therapy), AND BOTH of the following:
        - 1. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 2. The requested agent is in a different therapeutic class OR
      - c. The requested agent will be utilized for add-on therapy to existing dual therapy (triple therapy), AND ALL of the following:
        - 1. Patient is WHO functional class III or IV AND
        - 2. Patient has unacceptable or deteriorating clinical status despite established PAH pharmacotherapy AND
        - 3. All three agents in the triple therapy are from a different therapeutic class OR

459

- d. The requested agent will be utilized as part of triple therapy in a treatment naive patient AND BOTH of the following:
  - 1. Patient is classified as WHO functional class IV AND

2. The three agents being utilized consist of: ERA plus PDE5i plus prostanoid

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

## Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Drug is also subject to Part B versus Part D review.

Pulmonary Hypertension PA – Winrevair

# Drug Name(s)

Winrevair

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Pending CMS Review

Pyrimethamine PA

## Drug Name(s)

Daraprim

Pyrimethamine

## **Indications:**

All Medically-Accepted Indications.

## **Off-Label Uses:**

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

## Age Restriction:

**Prescriber Restrictions:** 

## **Coverage Duration:**

Approval will be for 6 months

Pyrukynd PA

## Drug Name(s)

Pyrukynd

Pyrukynd Taper Pack

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require the following:

1. Patient has a diagnosis of hemolytic anemia with pyruvate kinase deficiency (PKD)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of hemolytic anemia with pyruvate kinase deficiency (PKD) AND
- 3. Patient has had clinical benefit with the requested agent

## Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., hematologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Qbrexza PA

Drug Name(s)

Qbrexza

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has a diagnosis of primary axillary hyperhidrosis

Age Restriction:

Patient is within the FDA labeled age for the requested agent

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Quillichew PA

Drug Name(s)

Quillichew Er

Indications:

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Quillivant PA

Drug Name(s)

Quillivant Xr

Indications:

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Quinine PA

## Drug Name(s)

Qualaquin

Quinine Sulfate

#### Indications:

All Medically-Accepted Indications.

## Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Uncomplicated malaria OR
  - B. Babesiosis OR
  - C. An indication that is supported in CMS approved compendia for the requested agent

## Age Restriction:

## **Prescriber Restrictions:**

## **Coverage Duration:**

7 days for malaria, 10 days for babesiosis, 12 months for all other diagnoses

Qulipta PA

## Drug Name(s)

Qulipta

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of migraine AND
- 2. The requested agent is being used for migraine prophylaxis AND
- 3. Patient has 4 migraine or more headache days per month AND
- 4. Patient will NOT be using the requested agent in combination with another calcitonin gene-related peptide (CGRP) agent for migraine prophylaxis

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of migraine AND
- 3. The requested agent is being used for migraine prophylaxis AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with another calcitonin gene-related peptide (CGRP) agent for migraine prophylaxis

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Quviviq PA

Drug Name(s)

Quviviq

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Radicava PA

# Drug Name(s)

Edaravone

Radicava

Radicava Ors

Radicava Ors Starter Kit

#### Indications:

All FDA-Approved Indications.

#### **Off-Label Uses:**

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of amyotrophic lateral sclerosis (ALS) [also known as Lou Gehrig's disease] AND
- 2. ALL of the following:
  - A. Patient is able to perform most activities of daily living AND
  - B. Patient has had the diagnosis of amyotrophic lateral sclerosis (ALS) for a duration of 2 years or less AND
  - C. Patient has a baseline percent predicted forced vital capacity (FVC) or slow vital capacity (SVC) of 80% or greater AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of amyotrophic lateral sclerosis (ALS) [also known as Lou Gehrig's disease] AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be 6 months for initial, 12 months for renewal

Rasuvo PA

### Drug Name(s)

Rasuvo

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. Patient has tried and had an inadequate response to a generic methotrexate injectable agent OR
  - B. Patient has an intolerance or hypersensitivity to a generic methotrexate injectable agent OR
  - C. Patient has an FDA labeled contraindication to a generic methotrexate injectable agent OR
  - D. Prescriber has provided information that the patient has a physical or a mental disability that would prevent the patient from using a generic methotrexate injectable agent

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Rayos PA

# Drug Name(s)

Rayos

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - a. Patient has tried and failed generic oral prednisone AND at least 1 other different generic oral corticosteroid agent (e.g., dexamethasone, methylprednisolone) OR
  - b. Patient has an intolerance or hypersensitivity to generic oral prednisone AND at least 1 other different generic oral corticosteroid agent (e.g., dexamethasone, methylprednisolone) OR
  - c. Patient has an FDA labeled contraindication to generic oral prednisone AND at least 1 other different generic oral corticosteroid agent (e.g., dexamethasone, methylprednisolone)

**Age Restriction:** 

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 6 months

Reblozyl PA

# Drug Name(s)

Reblozyl

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. Patient has a diagnosis of Beta thalassemia requiring regular red blood cell (RBC) transfusions AND
    - ii. Patient's diagnosis was confirmed by BOTH of the following:
      - a. Hemoglobin analysis by hemoglobin electrophoresis or high-performance liquid chromatography AND
      - b. Genetic analysis for both Beta thalassemia and Alpha thalassemia mutations  $\ensuremath{\mathsf{OR}}$
  - B. Patient has a diagnosis of anemia associated with myelodysplastic syndrome with ring sideroblasts (MDS-RS) AND BOTH of the following:
    - i. Patient has very low-to-intermediate-risk disease AND
    - ii. BOTH of the following:
      - a. Patient has tried and had an inadequate response to an erythropoiesis stimulating agent (ESA) [e.g., Aranesp (darbepoetin alfa), Epogen/Procrit (epoetin alfa), Retacrit (epoetin alfa-epbx)] AND
      - b. Patient has required 2 or more red blood cell (RBC) units over 8 weeks OR
  - C. Patient has a diagnosis of anemia associated with myelodysplastic/myeloproliferative neoplasm with ring sideroblasts and thrombocytosis (MDS/MPN-RS-T) AND BOTH of the following:
    - i. Patient has very low-to-intermediate-risk disease AND
    - ii. BOTH of the following:
      - a. Patient has tried and had an inadequate response to an erythropoiesis stimulating agent (ESA) [e.g., Aranesp (darbepoetin alfa), Epogen/Procrit (epoetin alfa), Retacrit (epoetin alfa-epbx)] AND
      - b. Patient has required 2 or more red blood cell (RBC) units over 8 weeks OR
  - D. Patient has a diagnosis of anemia associated with myelodysplastic syndromes (MDS) without previous erythropoiesis stimulating agent use (ESA-naive) AND BOTH of the following:
    - i. Patient has very low-to-intermediate-risk disease AND
    - ii. Patient has required 2 or more red blood cell (RBC) units over 8 weeks AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

**Prescriber Restrictions:** 

Prescriber is a specialist in the area of the patient's diagnosis (e.g., hematologist, oncologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Initial approval: 15 wks for Beta thalassemia, 12 mos for all other diagnoses. 12 mos for renewal.

### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of Beta thalassemia requiring regular red blood cell (RBC) transfusions OR
  - B. Patient has a diagnosis of anemia associated with myelodysplastic syndrome with ring sideroblasts (MDS-RS) OR
  - C. Patient has a diagnosis of anemia associated with myelodysplastic/myeloproliferative neoplasm with ring sideroblasts and thrombocytosis (MDS/MPN-RS-T) OR
  - D. Patient has a diagnosis of anemia associated with myelodysplastic syndromes (MDS) without previous erythropoiesis stimulating agent use (ESA-naive) AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Recorlev PA

# Drug Name(s)

Recorlev

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of endogenous hypercortisolemia with Cushing's syndrome AND
- 2. ONE of the following:
  - A. Patient had an inadequate response to pituitary surgical resection OR
  - B. Patient is NOT a candidate for pituitary surgical resection AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to pasireotide OR
  - B. Patient has an intolerance or hypersensitivity to pasireotide OR
  - C. Patient has an FDA labeled contraindication to pasireotide

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of endogenous hypercortisolemia with Cushing's syndrome AND
- 3. Patient has had clinical benefit with the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Regranex PA

# Drug Name(s)

Regranex

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. Patient has a diagnosis of lower extremity diabetic neuropathic ulcer(s) that extends into the subcutaneous tissue or beyond AND
    - ii. The ulcer(s) intended for treatment has an adequate blood supply OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Relexxii PA

# Drug Name(s)

Methylphenidate Hcl Er (Relexxii)

Relexxii

### Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Relistor Inj PA

Drug Name(s)

Relistor Injection

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. ONE of the following diagnoses:
  - A. Patient has opioid-induced constipation (OIC) with advanced illness or pain caused by active cancer and is receiving palliative care OR
  - B. Patient has opioid induced constipation (OIC) and chronic non-cancer pain, including patients with chronic pain related to prior cancer or its treatment AND
- 2. Patient has chronic use of an opioid agent in the past 90 days AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to lactulose OR
  - B. Patient has an intolerance or hypersensitivity to lactulose OR
  - C. Patient has an FDA labeled contraindication to lactulose

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Relistor Tablet PA

# Drug Name(s)

**Relistor Tablet** 

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has opioid-induced constipation (OIC) and chronic non-cancer pain, including patients with chronic pain related to prior cancer or its treatment AND
- 2. Patient has chronic use of an opioid agent in the past 90 days AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to lactulose OR
  - B. Patient has an intolerance or hypersensitivity to lactulose OR
  - C. Patient has an FDA labeled contraindication to lactulose

# Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 months

Repatha PA

Drug Name(s)

Repatha

Repatha Pushtronex System

Repatha Sureclick

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Reyvow PA

# Drug Name(s)

Reyvow

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. The requested agent will be used for the treatment of acute migraine with or without aura AND
- 2. ONE of the following:
  - A. Patient has tried and had an inadequate response to TWO triptan agents with differing active ingredients (e.g., sumatriptan, rizatriptan) OR
  - B. Patient has an intolerance or hypersensitivity to TWO triptan agents with differing active ingredients OR
  - C. Patient has an FDA labeled contraindication to TWO triptan agents with differing active ingredients AND
- 3. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, ergotamine, acute CGRP)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. The requested agent will be used for the treatment of acute migraine with or without aura AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, ergotamine, acute CGRP)

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Rezdiffra PA

Drug Name(s)

Rezdiffra

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

Decompensated cirrhosis AND

Moderate to severe hepatic impairment (Child-Pugh Class B or C)

**Required Medical Information:** 

Pending CMS Review

Rezurock PA

# Drug Name(s)

Rezurock

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of chronic graft-versus-host disease (chronic GVHD) AND
- 2. Patient has failed at least two prior lines of systemic therapy

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of chronic graft-versus-host disease (chronic GVHD) AND
- 3. Patient has had clinical benefit with the requested agent

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., hematologist, oncologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Ritalin LA PA

# Drug Name(s)

Methylphenidate Hcl Er (Ritalin LA)

Ritalin La

### Indications:

All FDA-Approved Indications.

**Off-Label Uses:** 

# **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Ritalin PA

# Drug Name(s)

Methylphenidate Hcl (Ritalin)

Ritalin

### Indications:

All FDA-Approved Indications.

**Off-Label Uses:** 

# **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Rivfloza PA

# Drug Name(s)

Rivfloza

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of primary hyperoxaluria type 1 (PH1) confirmed by ONE of the following:
  - A. Genetic testing of the AGXT gene indicates a pathogenic mutation OR
  - B. Liver biopsy demonstrates absent or significantly reduced alanine:glyoxylate aminotransferase (AGT) activity AND
- 2. The requested agent will be used to lower urinary oxalate levels AND
- 3. Patient has an estimated GFR (eGFR) greater than or equal to 30 mL/min/1.73^2 AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of primary hyperoxaluria type 1 (PH1) AND
- 3. The requested agent will be used to lower urinary oxalate levels AND
- 4. Patient has an estimated GFR (eGFR) greater than or equal to 30 mL/min/1.73^2 AND
- 5. Patient has had clinical benefit with the requested agent AND
- 6. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, nephrologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be 6 months for initial, 12 months for renewal

Roflumilast PA

# Drug Name(s)

Daliresp

Roflumilast

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication for the requested agent AND
- 2. ONE of the following:
  - A. Patient has tried and had an inadequate response to an agent from TWO of the following categories:
    - i. long-acting beta-2 agonist (LABA) [e.g., salmeterol]
    - ii. long-acting muscarinic antagonist (LAMA) [e.g., umeclidinium]
    - iii. inhaled corticosteroid (ICS) [e.g., fluticasone] OR
  - B. Patient has an intolerance or hypersensitivity to an agent from TWO of the following categories:
    - i. long-acting beta-2 agonist (LABA) [e.g., salmeterol]
    - ii. long-acting muscarinic antagonist (LAMA) [e.g., umeclidinium]
    - iii. inhaled corticosteroid (ICS) [e.g., fluticasone] OR
  - C. Patient has an FDA labeled contraindication to an agent from TWO of the following categories:
    - i. long-acting beta-2 agonist (LABA) [e.g., salmeterol]
    - ii. long-acting muscarinic antagonist (LAMA) [e.g., umeclidinium]
    - iii. inhaled corticosteroid (ICS) [e.g., fluticasone]

# Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Rybelsus PA

# Drug Name(s)

Rybelsus

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

Requested agent will be used for weight loss alone

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of type 2 diabetes mellitus AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 180 days OR
  - C. ALL of the following:
    - i. ONE of the following:
      - 1. Patient's medication history includes use of a non glucagon-like peptide-1 (GLP-1) oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) within the past 90 days OR
      - 2. Patient had an ineffective treatment response to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 3. Patient has an intolerance or hypersensitivity to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 4. Patient has an FDA labeled contraindication to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent AND iii. Patient will NOT be using the requested agent in combination with another GLP-1 agonist agent, or an agent containing a GLP-1 agonist AND
    - iv. Patient will NOT be using the requested agent in combination with an agent containing a dipeptidyl peptidase-4 (DPP-4) inhibitor

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Samsca PA

### Drug Name(s)

Samsca

Tolvaptan

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

Any underlying liver disease, including cirrhosis AND FDA labeled contraindications to the request agent **Required Medical Information:** 

Criteria for approval require ALL of the following:

- 1. The requested agent was initiated (or re-initiated) in the hospital AND
- 2. Prior to initiating the requested agent, the patient has or had a diagnosis of clinically significant hypervolemic or euvolemic hyponatremia defined by ONE of the following:
  - A. Serum sodium is less than 125 mEq/L OR
  - B. Serum sodium is 125 mEq/L or greater AND patient has symptomatic hyponatremia that has resisted correction with fluid restriction AND
- 3. Medications known to cause hyponatremia have been evaluated and discontinued when appropriate AND
- 4. Patient has NOT already received 30 days of therapy with the requested agent following the most recent hospitalization for initiation of therapy AND
- 5. The requested dose is within the FDA labeled dosing for the requested indication (Recommended starting dose is 15 mg once daily. Dosage may be increased at intervals greater than or equal to 24 hours to 30 mg once daily, and to a maximum of 60 mg once daily as needed to raise serum sodium. Do not administer for more than 30 days to minimize the risk of liver injury.)

# Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 30 days

Sapropterin PA

# Drug Name(s)

Javygtor

Kuvan

Sapropterin Dihydrochloride

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of phenylketonuria (PKU) AND
- 2. Prescriber has submitted a baseline blood Phe level measured prior to initiation of therapy with the requested agent, which is above the recommended levels indicated for the patient's age range or condition AND
- 3. Patient will NOT be using the requested agent in combination with Palynziq (pegvaliase-pqpz) for the requested indication AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of phenylketonuria (PKU) AND
- 3. ONE of the following:
  - a. Patient's blood Phe levels are being maintained within the acceptable range OR
  - b. Patient has had a decrease in blood Phe level from baseline AND
- 4. Patient will NOT be using the requested agent in combination with Palynziq (pegvaliase-pqpz) for the requested indication AND
- 5. The requested dose is within FDA labeled dosing for the requested indication

### Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., metabolic or genetic disorders) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Initial: 2 months if dose is 5 to less than 20 mg/kg/day, 1 month if 20 mg/kg/day Renewal: 12 months Other Criteria:

Self - Administered Oncology PA

# Drug Name(s)

Abiraterone Acetate

Afinitor

Afinitor Disperz

Akeega

Alecensa

Alunbrig

Augtyro

Ayvakit

Balversa

Besremi

Bexarotene Capsule

Bosulif

Braftovi

Brukinsa

Cabometyx

Calquence

Caprelsa

Cometriq

Copiktra

Cotellic

Daurismo

Erivedge

Erleada

Erlotinib Hcl

Everolimus

Exkivity

Fotivda

Fruzagla

Gavreto

Gefitinib

Gilotrif

.

Gleevec

Ibrance Iclusig

Idhifa

Imatinib Mesylate

Imbruvica

Inlyta

Inqovi

Inrebic

Iressa

Iwilfin

Jakafi

Jaypirca

Kisqali

Kisqali Femara 200 Dose

Kisqali Femara 400 Dose

Kisqali Femara 600 Dose

Koselugo

Krazati

Lapatinib Ditosylate

Lazcluze

Lenalidomide

Lenvima 10 Mg Daily Dose

Lenvima 12Mg Daily Dose

Lenvima 14 Mg Daily Dose

Lenvima 18 Mg Daily Dose

Lenvima 20 Mg Daily Dose

Lenvima 24 Mg Daily Dose

Lenvima 4 Mg Daily Dose

Lenvima 8 Mg Daily Dose

Lonsurf

Lorbrena

Lumakras

Lynparza

Lytgobi

Matulane

Mekinist

Mektovi

Nerlynx

Nexavar

Ninlaro

Nubega

Odomzo

Ojemda

Ogsiveo

0631166

Ojjaara

Onureg

Orgovyx

Orserdu

Pazopanib Hcl

Pemazyre

Pigray 200Mg Daily Dose

Piqray 250Mg Daily Dose

Pigray 300Mg Daily Dose

Pomalyst

Qinlock

Retevmo

Revlimid

Rezlidhia

Rozlytrek

Rubraca

Rydapt

Scemblix

Sorafenib

Sprycel

Stivarga

Sunitinib Malate

Sutent

Tabrecta

Tafinlar

Tagrisso

Talzenna

Targretin Capsule

Tasigna

Tazverik

Tepmetko

Thalomid

Tibsovo

Torpenz

Tretinoin Capsule 10Mg

Truqap

Tukysa

Turalio

Tykerb

Vanflyta

Venclexta

Venclexta Starting Pack

Verzenio

Vitrakvi

Vizimpro

Vonjo

Voranigo

Votrient

Welireg

Xalkori

Xospata

Xpovio

Xtandi

Yonsa

Zejula

Zelboraf

Zolinza

Zydelig

Zykadia

Zytiga

#### Indications:

All Medically-Accepted Indications.

#### **Off-Label Uses:**

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. ALL of the following:
    - i. Genetic testing has been completed, if required, for therapy with the requested agent and results indicate the requested agent is appropriate AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent AND iii. ONE of the following:
      - a. The requested agent is FDA labeled or supported by CMS approved compendia as a first-line therapy for the requested indication OR
      - b. Patient has tried appropriate FDA labeled or CMS approved compendia supported therapy that are indicated as first-line therapy for the requested indication OR
      - c. Patient has an intolerance or hypersensitivity to the first-line therapy for the requested indication OR
      - d. Patient has an FDA labeled contraindication to the first-line therapy for the requested indication AND
    - iv. Patient does NOT have any FDA labeled limitations of use that is not otherwise supported in NCCN guidelines AND

Criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

v. ONE of the following:

- a. The requested agent is not Bosulif OR
- b. The requested agent is Bosulif AND ONE of the following:
  - 1. Patient's medication history indicates use of imatinib OR dasatinib for the requested indication (if applicable) OR
  - 2. Patient has an intolerance or hypersensitivity to imatinib OR dasatinib OR
  - 3. Patient has an FDA labeled contraindication to imatinib OR dasatinib OR
  - 4. CMS approved compendia does not support the use of imatinib OR dasatinib for the requested indication OR
  - 5. Prescriber has provided information in support of use of Bosulif over imatinib OR dasatinib for the requested indication AND

# vi. ONE of the following:

- a. The requested agent is not Calquence OR
- b. The requested agent is Calquence AND ONE of the following:
  - 1. Patient's medication history indicates use of Brukinsa OR Imbruvica for the requested indication (if applicable) OR
  - 2. Patient has an intolerance or hypersensitivity to Brukinsa OR Imbruvica OR
  - 3. Patient has an FDA labeled contraindication to Brukinsa OR Imbruvica OR
  - 4. CMS approved compendia do not support the use of Brukinsa OR Imbruvica for the requested indication OR
  - 5. Prescriber has provided information in support of use of Calquence over Brukinsa OR Imbruvica for the requested indication

Signifor LAR PA

### Drug Name(s)

Signifor Lar

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

Severe hepatic impairment (i.e., Child Pugh C)

# **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of acromegaly AND ONE of the following:
    - i. Patient had an inadequate response to surgery as indicated by growth hormone and serum IGF-1 levels that are above the reference ranges for the patient's gender and age OR
    - ii. Patient is NOT a candidate for surgery OR
  - B. Patient has a diagnosis of Cushing's disease (CD) AND ONE of the following:
    - i. Patient had an inadequate response to pituitary surgical resection OR
    - ii. Patient is NOT a candidate for pituitary surgical resection OR
  - C. Patient has an indication that is supported in CMS approved compendia for the requested agent

Criteria for renewal approval require BOTH of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of acromegaly AND ONE of the following:
    - i. Patient has growth hormone and serum IGF-1 levels that are within normal limits for patient's gender and age reference range OR
    - ii. Patient has had clinical improvement (e.g., reduction in tumor size, decreased headaches, improved cardiovascular or respiratory symptoms) OR
  - B. Patient has a diagnosis of Cushing's disease (CD) AND BOTH of the following:
    - i. Patient has a urinary free cortisol level less than or equal to the upper limit of normal AND
    - ii. Patient has had improvement in at least ONE of the following clinical signs and symptoms:
      - 1. Fasting plasma glucose OR
      - 2. Hemoglobin A1c OR
      - 3. Hypertension OR
      - 4. Weight OR
  - C. BOTH of the following:
    - i. Patient has an indication that is supported in CMS approved compendia for the requested agent AND

ii. Patient has had clinical benefit with the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Initial: Acromegaly - 6 months, CD - 7 months, All other diagnoses - 12 months, Renewal: 12 months

Signifor PA

# Drug Name(s)

Signifor

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

Severe hepatic impairment (i.e., Child Pugh C)

# **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of Cushing's disease (CD) AND ONE of the following:
    - i. Patient had an inadequate response to pituitary surgical resection OR
    - ii. Patient is NOT a candidate for pituitary surgical resection OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

Criteria for renewal approval require BOTH of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of Cushing's disease (CD) AND BOTH of the following:
    - i. Patient has a urinary free cortisol level less than or equal to the upper limit of normal AND
    - ii. Patient has had improvement in at least ONE of the following clinical signs and symptoms:
      - 1. Fasting plasma glucose OR
      - 2. Hemoglobin A1c OR
      - 3. Hypertension OR
      - 4. Weight OR
  - B. BOTH of the following:
    - i. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient has had clinical benefit with the requested agent

### **Age Restriction:**

### **Prescriber Restrictions:**

### **Coverage Duration:**

Initial approval: 6 months for CD, 12 months for all other diagnoses, Renewal approval: 12 months Other Criteria:

Sivextro PA

### Drug Name(s)

Sivextro

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has ONE of the following:
  - A. BOTH of the following:
    - i. A documented acute bacterial skin and skin structure infection (ABSSSI) defined as a bacterial infection of the skin with a lesion size area of at least 75 cm2 (lesion size measured by the area of redness, edema, or induration) AND
    - ii. The infection is due to Staphylococcus aureus, Streptococcus pyogenes, Streptococcus agalactiae, Streptococcus anginosus, Streptococcus intermedius, Streptococcus constellatus, or Enterococcus faecalis OR
  - B. Another indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. The requested agent is prescribed by an infectious disease specialist or the prescriber has consulted with an infectious disease specialist on treatment of this patient OR
  - B. The requested agent is NOT prescribed by an infectious disease specialist or the prescriber has NOT consulted with an infectious disease specialist on treatment of this patient AND ONE of the following:
    - i. There is documentation of resistance to TWO of the following: beta-lactams, macrolides, clindamycin, tetracycline, or co-trimoxazole at the site of infection OR
    - ii. Patient has an intolerance or hypersensitivity to TWO of the following: beta-lactams, macrolides, clindamycin, tetracyclines, or co-trimoxazole OR
    - iii. Patient has an FDA labeled contraindication to TWO of the following: beta-lactams, macrolides, clindamycin, tetracyclines, or co-trimoxazole OR
    - iv. There is documentation of resistance to vancomycin at the site of infection OR
    - v. Patient has an intolerance or hypersensitivity to vancomycin OR
    - vi. Patient has an FDA labeled contraindication to vancomycin AND
- 3. Patient will NOT be using the requested agent in combination with linezolid for the same infection AND
- 4. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

#### Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be 6 days for ABSSSI or 30 days for all other indications

Skyclarys PA

# Drug Name(s)

Skyclarys

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for initial approval require the following:

1. Patient has a diagnosis of Friedreich's ataxia (FA, FRDA) with genetic analysis confirming mutation in the frataxin (FXN) gene

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Friedreich's ataxia (FA, FRDA) AND
- 3. Patient has had clinical benefit with the requested agent

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cardiologist, geneticist, neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Sodium Oxybate PA

# Drug Name(s)

Sodium Oxybate

**Xyrem** 

#### **Indications:**

All Medically-Accepted Indications.

### **Off-Label Uses:**

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of narcolepsy with cataplexy OR
  - B. Patient has a diagnosis of narcolepsy with excessive daytime sleepiness AND BOTH of the following:
    - i. ONE of the following:
      - a. Patient is under 18 years of age OR
      - b. ONE of the following:
        - 1. Patient has tried and had an inadequate response to modafinil or armodafinil OR
        - 2. Patient has an intolerance or hypersensitivity to modafinil or armodafinil OR
        - 3. Patient has an FDA labeled contraindication to modafinil or armodafinil AND
    - ii. ONE of the following:
      - a. Patient has tried and had an inadequate response to ONE standard stimulant agent (e.g., methylphenidate) OR
      - b. Patient has an intolerance or hypersensitivity to ONE standard stimulant agent (e.g., methylphenidate) OR
      - c. Patient has an FDA labeled contraindication to ONE standard stimulant agent (e.g., methylphenidate) OR
  - C. Patient has another indication that is supported in CMS approved compendia for the requested agent

### Age Restriction:

Patient is 7 years of age or over

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Somatostatin Analogs PA - Lanreotide

#### Drug Name(s)

Somatuline Depot

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent OR
  - B. ONE of the following:
    - i. Patient has a diagnosis of acromegaly AND ONE of the following:
      - a. Patient is not a candidate for surgical resection or pituitary radiation therapy OR
      - b. The requested agent is for adjunctive therapy with pituitary radiation therapy OR
      - c. Patient had an inadequate response to surgery or pituitary radiation therapy as indicated by growth hormone levels or serum IGF-1 levels that are above the reference range OR
    - ii. Patient has a diagnosis of gastroenteropancreatic neuroendocrine tumors AND BOTH of the following:
      - a. The tumors are well or moderately differentiated AND
      - b. ONE of the following:
        - 1. The tumors are unresectable, locally advanced OR
        - 2. Patient has metastatic disease OR
    - iii. Patient has a diagnosis of carcinoid syndrome OR
    - iv. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

### Age Restriction:

#### **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be 6 months for initial, 12 months for renewal

#### Other Criteria:

Criteria for renewal approval require ALL of the following:

1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND

- 2. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 3. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. BOTH of the following:
    - i. ONE of the following:
      - 1. Patient has a diagnosis of acromegaly OR
      - 2. Patient has a diagnosis of metastatic OR unresectable, locally advanced, well or moderately differentiated gastroenteropancreatic neuroendocrine tumors OR
      - 3. Patient has a diagnosis of carcinoid syndrome OR
      - 4. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Somatostatin Analogs PA – Mycapssa

#### Drug Name(s)

Mycapssa

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - B. ONE of the following:
    - i. BOTH of the following:
      - a. Patient has a diagnosis of acromegaly AND
      - b. Patient has responded to and tolerated treatment with octreotide or lanreotide OR
    - ii. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of acromegaly OR
  - B. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

#### Age Restriction:

**Prescriber Restrictions:** 

## **Coverage Duration:**

Approval will be 9 months for initial, 12 months for renewal

Somatostatin Analogs PA – Octreotide

#### Drug Name(s)

Octreotide Acetate

Sandostatin

#### **Indications:**

All Medically-Accepted Indications.

#### **Off-Label Uses:**

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - B. ONE of the following:
    - i. Patient has a diagnosis of acromegaly AND ONE of the following:
      - a. Patient is not a candidate for surgical resection or pituitary radiation therapy OR
      - b. The requested agent is for adjunctive therapy with pituitary radiation therapy OR
      - c. Patient had an inadequate response to surgery or pituitary radiation therapy as indicated by growth hormone levels or serum IGF-1 levels that are above the reference range OR
    - ii. Patient has severe diarrhea and/or flushing episodes associated with metastatic carcinoid tumors OR
    - iii. Patient has profuse watery diarrhea associated with Vasoactive Intestinal Peptide (VIP) secreting tumors OR
    - iv. Patient has a diagnosis of dumping syndrome AND ONE of the following:
      - a. Patient has tried and had an inadequate response to acarbose OR
      - b. Patient has an intolerance or hypersensitivity to acarbose OR
      - c. Patient has an FDA labeled contraindication to acarbose OR
    - v. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
- 2. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

### Age Restriction:

#### **Prescriber Restrictions:**

### **Coverage Duration:**

Approval will be 6 months for initial, 12 months for renewal

#### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of acromegaly OR
  - B. Patient has severe diarrhea and/or flushing episodes associated with metastatic carcinoid tumors OR
  - C. Patient has profuse watery diarrhea associated with Vasoactive Intestinal Peptide (VIP) secreting tumors OR
  - D. Patient has a diagnosis of dumping syndrome OR
  - E. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Somatostatin Analogs PA - Sandostatin LAR

### Drug Name(s)

Sandostatin Lar Depot

Indications:

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. ONE of the following:
  - A. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND ONE of the following:
    - i. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
    - ii. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR
  - B. ONE of the following:
    - i. Patient has a diagnosis of acromegaly AND ONE of the following:
      - a. Patient is not a candidate for surgical resection or pituitary radiation therapy OR
      - b. The requested agent is for adjunctive therapy with pituitary radiation therapy OR
      - c. Patient had an inadequate response to surgery or pituitary radiation therapy as indicated by growth hormone levels or serum IGF-1 levels that are above the reference range OR
    - ii. Patient has severe diarrhea and/or flushing episodes associated with metastatic carcinoid tumors OR
    - iii. Patient has profuse watery diarrhea associated with Vasoactive Intestinal Peptide (VIP) secreting tumors OR
    - iv. Patient has a diagnosis of dumping syndrome AND ONE of the following:
      - a. Patient has tried and had an inadequate response to acarbose OR
      - b. Patient has an intolerance or hypersensitivity to acarbose OR
      - c. Patient has an FDA labeled contraindication to acarbose OR
    - v. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
- 2. Patient has responded to and tolerated octreotide for a minimum of 2 weeks prior to starting therapy with Sandostatin LAR AND
- 3. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be 6 months for initial, 12 months for renewal

# Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of acromegaly OR
  - B. Patient has severe diarrhea and/or flushing episodes associated with metastatic carcinoid tumors OR
  - C. Patient has profuse watery diarrhea associated with Vasoactive Intestinal Peptide (VIP) secreting tumors OR
  - D. Patient has a diagnosis of dumping syndrome OR
  - E. Patient has another indication that is supported in CMS approved compendia for the requested agent AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

Somatostatin Analogs PA – Somavert

#### Drug Name(s)

Somavert

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of acromegaly AND ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent AND provided clinical justification to support that the patient is at risk if therapy is changed OR C. BOTH of the following:
    - i. ONE of the following:
      - a. Patient is not a candidate for surgical resection or pituitary radiation therapy OR
      - b. The requested agent is for adjunctive therapy with pituitary radiation therapy OR
      - c. Patient had an inadequate response to surgery or pituitary radiation therapy as indicated by serum IGF-1 levels that are above the reference range AND
    - ii. ONE of the following:
      - a. Patient has tried and had an inadequate response to octreotide or Somatuline Depot (lanreotide) OR
      - b. Patient has an intolerance or hypersensitivity to octreotide or Somatuline Depot (lanreotide) OR
      - c. Patient has an FDA labeled contraindication to octreotide or Somatuline Depot (lanreotide) AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of acromegaly AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be 6 months for initial, 12 months for renewal

Sovaldi PA

### Drug Name(s)

Sovaldi

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of hepatitis C confirmed by serological markers AND
- 2. Prescriber has screened the patient for current or prior hepatitis B viral (HBV) infection and if positive, will monitor the patient for HBV flare-up or reactivation during and after treatment with the requested agent AND
- 3. The requested agent will be used in a treatment regimen and length of therapy that is supported in FDA approved labeling or AASLD/IDSA guidelines for the patient's diagnosis and genotype AND
- 4. The requested dose is within FDA labeled dosing or supported in AASLD/IDSA guideline dosing for the requested indication AND
- 5. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
  - C. Patient has an FDA labeled contraindication or hypersensitivity to the preferred agent: Mavyret for supported genotypes OR
  - D. Prescriber has provided information based on FDA approved labeling or AASLD/IDSA guidelines supporting the use of the non-preferred agent for the patient's diagnosis and genotype over the preferred agent: Mavyret for supported genotypes

#### Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, hepatologist or infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Duration of therapy: Based on FDA approved labeling or AASLD/IDSA guideline supported **Other Criteria**:

Spevigo SC PA

Drug Name(s)

Spevigo

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Strensiq PA

### Drug Name(s)

Strensiq

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Perinatal or infantile-onset hypophosphatasia OR
  - B. Juvenile-onset hypophosphatasia AND
- 2. Patient has documentation (i.e., medical records) of clinical manifestations to support the diagnosis of hypophosphatasia at the age of onset prior to age 18 (e.g., vitamin B6-dependent seizures, skeletal abnormalities such as rachitic chest deformity leading to respiratory problems or bowed arms/legs, "failure to thrive") AND
- 3. Patient has documentation (i.e., medical records) of radiographic imaging to support the diagnosis of hypophosphatasia at the age of onset prior to age 18 (e.g., infantile rickets, alveolar bone loss, craniosynostosis, fractures) AND
- 4. Patient has documentation (i.e., medical records) of confirmed mutation(s) in the ALPL gene that encodes the tissue non-specific isoenzyme of alkaline phosphatase (TNSALP) AND
- 5. Patient has documentation (i.e., medical records) of a measured total serum alkaline phosphatase (ALP) level that is below the normal lab reference range for age and sex AND
- 6. Patient has documentation (i.e., medical records) of ONE of the following:
  - A. Elevated urine concentration of phosphoethanolamine (PEA) OR
  - B. Elevated serum concentration of pyridoxal 5'-phosphate (PLP) in the absence of vitamin supplements within one week prior to the test OR
  - C. Elevated urinary inorganic pyrophosphate (PPi) AND
- 7. The requested dose is within FDA labeled dosing (based on the patient's weight) for the requested indication

#### Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., endocrinologist or geneticist with expertise in metabolic bone diseases) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has ONE of the following diagnoses:
  - A. Perinatal or infantile-onset hypophosphatasia OR

- B. Juvenile-onset hypophosphatasia AND
- 3. There is documentation (i.e., medical records) that the patient has had a decrease from baseline (before treatment with the requested agent) in at least ONE of the following levels:
  - A. Urine concentration of phosphoethanolamine (PEA) OR
  - B. Serum concentration of pyridoxal 5'-phosphate (PLP) in the absence of vitamin supplements within one week prior to the test OR
  - C. Urinary inorganic pyrophosphate (PPi) AND
- 4. Patient has documentation (i.e., medical records) of clinical improvement and/or stabilization with the requested agent (e.g., improvement in respiratory status, growth, pain, radiographic findings, other symptoms associated with the disease) AND
- 5. The requested dose is within FDA labeled dosing (based on the patient's weight) for the requested indication

Substrate Reduction Therapy PA – Cerdelga

#### Drug Name(s)

Cerdelga

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of Gaucher disease type 1 (GD1) confirmed by ONE of the following:
- A. A baseline (prior to therapy for the requested indication) glucocerebrosidase enzyme activity of less than or equal to 15% of mean normal in peripheral blood leukocytes, fibroblasts, or other nucleated cells OR
- B. Confirmation of genetic mutation of the glucocerebrosidase (GBA) gene with two disease-causing alleles AND
- 2. Patient is a CYP2D6 extensive metabolizer (EMs), intermediate metabolizer (IMs), or poor metabolizer (PMs) established by a genetic test AND
- 3. Prescriber has drawn baseline (prior to therapy for the requested indication) measurements of hemoglobin level, platelet count, liver volume, and spleen volume AND
- 4. Prior to any treatment for the intended diagnosis, the patient has had at least ONE of the following clinical presentations:
- A. Anemia [defined as mean hemoglobin (Hb) level below the testing laboratory's lower limit of the normal range based on age and gender] OR
- B. Thrombocytopenia (defined as platelet count of less than 100,000 per microliter) OR
- C. Hepatomegaly OR
- D. Splenomegaly OR
- E. Growth failure (i.e., growth velocity is below the standard mean for age) OR
- F. Evidence of bone disease with other causes ruled out

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Gaucher disease type 1 (GD1) AND
- 3. Patient has had improvements or stabilization with the requested agent as indicated by ONE of the following:
- A. Spleen volume OR
- B. Hemoglobin level OR
- C. Liver volume OR
- D. Platelet count OR
- E. Growth OR
- F. Bone pain or crisis

#### Age Restriction:

# **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., endocrinologist, geneticist, hematologist, specialist in metabolic diseases) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

 ${\bf Substrate\ Reduction\ The rapy\ PA-Miglustat}$ 

Drug Name(s)

Miglustat

Yargesa

Zavesca

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Sucraid PA

### Drug Name(s)

Sucraid

#### **Indications:**

All FDA-Approved Indications.

#### **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of congenital sucrase-isomaltase deficiency (CSID) AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of congenital sucrase-isomaltase deficiency (CSID) AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, geneticist, endocrinologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

#### **Coverage Duration:**

Approval will be for 12 months

Sunosi PA

#### Drug Name(s)

Sunosi

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of excessive daytime sleepiness associated with narcolepsy AND BOTH of the following:
    - i. ONE of the following:
      - 1. Patient has tried and had an inadequate response to modafinil or armodafinil OR
      - 2. Patient has an intolerance or hypersensitivity to modafinil or armodafinil OR
      - 3. Patient has an FDA labeled contraindication to modafinil or armodafinil AND
    - ii. ONE of the following:
      - 1. Patient has tried and had an inadequate response to ONE standard stimulant agent (e.g., methylphenidate) OR
      - 2. Patient has an intolerance or hypersensitivity to ONE standard stimulant agent (e.g., methylphenidate) OR
      - 3. Patient has an FDA labeled contraindication to ONE standard stimulant agent (e.g., methylphenidate) OR
  - B. Patient has a diagnosis of excessive daytime sleepiness associated with obstructive sleep apnea (OSA) AND ONE of the following:
    - i. Patient has tried and had an inadequate response to modafinil or armodafinil OR
    - ii. Patient has an intolerance or hypersensitivity to modafinil or armodafinil OR
    - iii. Patient has an FDA labeled contraindication to modafinil or armodafinil AND
- 2. Patient will NOT be using the requested agent in combination with modafinil, armodafinil, or a standard stimulant agent (e.g., methylphenidate) for the requested indication

#### Age Restriction:

Patient is 18 years of age or over

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Approval will be for 12 months

#### Other Criteria:

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of excessive daytime sleepiness associated with narcolepsy OR

- B. Patient has a diagnosis of excessive daytime sleepiness associated with obstructive sleep apnea (OSA) AND
- 3. Patient will NOT be using the requested agent in combination with modafinil, armodafinil, or a standard stimulant agent (e.g., methylphenidate) for the requested indication AND
- 4. Patient has had clinical benefit with the requested agent

Symdeko PA

### Drug Name(s)

Symdeko

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of cystic fibrosis AND
- 2. ONE of the following:
  - A. Patient has the presence of the F508del mutation on both alleles (homozygous) of the CFTR gene confirmed by genetic testing OR
  - B. Patient has ONE of the CFTR gene mutations or a mutation in the CFTR gene that is responsive based on in vitro data, as indicated in the FDA label, confirmed by genetic testing OR
  - C. Patient has another CFTR gene mutation(s) that is responsive to the requested agent, as indicated in the FDA label, confirmed by genetic testing AND
- 3. Patient will NOT be using the requested agent in combination with another CFTR modulator agent for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of cystic fibrosis AND
- 3. Patient has had improvement or stabilization with the requested agent [e.g., improvement in FEV1 from baseline, increase in weight/BMI, improvement from baseline Cystic Fibrosis Questionnaire-Revised (CFQ-R) Respiratory Domain score, improvements in respiratory symptoms (cough, sputum production, and difficulty breathing), and/or reduced number of pulmonary exacerbations] AND
- 4. Patient will NOT be using the requested agent in combination with another CFTR modulator agent for the requested indication

#### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cystic fibrosis, pulmonologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Symproic PA

### Drug Name(s)

Symproic

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has opioid-induced constipation (OIC) and chronic non-cancer pain, including patients with chronic pain related to prior cancer or its treatment AND
- 2. Patient has chronic use of an opioid agent in the past 90 days AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to lactulose OR
  - B. Patient has an intolerance or hypersensitivity to lactulose OR
  - C. Patient has an FDA labeled contraindication to lactulose

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Tarpeyo PA

### Drug Name(s)

Tarpeyo

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of primary immunoglobulin A nephropathy (IgAN) confirmed by kidney biopsy AND
- 2. The requested agent will be used to reduce the loss of kidney function AND
- 3. Patient is at risk of disease progression as shown by ONE of the following:
  - A. A urine protein-to-creatinine ratio (UPCR) greater than or equal to 0.8 g/g OR
  - B. Proteinuria greater than or equal to 1 g/day AND
- 4. ONE of the following:
  - A. Patient is currently being treated with an ACEI or ARB (e.g., benazepril, lisinopril, losartan), or a combination medication containing an ACE or ARB OR
  - B. Patient has an intolerance or hypersensitivity to an ACEI or ARB, or a combination medication containing an ACE or ARB OR
  - C. Patient has an FDA labeled contraindication to an ACEI or ARB, or a combination medication containing an ACE or ARB AND
- 5. Patient has not previously been treated with a course of therapy (10 months) with the requested agent AND
- 6. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., nephrologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

No prior Tarpeyo use, approve 10 months. Prior use - see Other Criteria.

#### Other Criteria:

Prior Tarpeyo use, approve remainder of 10 months total course of therapy.

Tasimelteon Capsule PA

## Drug Name(s)

Hetlioz

Tasimelteon

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. Patient has a diagnosis of Non-24-hour sleep-wake disorder AND
    - ii. Patient is totally blind (i.e., no light perception) OR
  - B. BOTH of the following:
    - i. Patient has a diagnosis of Smith-Magenis Syndrome (SMS) confirmed by the presence of ONE of the following genetic mutations:
      - A. A heterozygous deletion of 17p11.2 OR
      - B. A heterozygous pathogenic variant involving RAI1 AND
    - ii. The requested agent is being used to treat nighttime sleep disturbances associated with SMS

#### Age Restriction:

For diagnosis of Non-24-hour sleep-wake disorder, patient is 18 years of age or over. For diagnosis of Smith-Magenis Syndrome (SMS), patient is 16 years of age or over.

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist, sleep specialist, psychiatrist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Tavalisse PA

# Drug Name(s)

Tavalisse

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of chronic immune (idiopathic) thrombocytopenia (ITP) AND
- 2. ONE of the following:
  - A. Patient has tried and had an insufficient response to a corticosteroid, another thrombopoietin receptor agonist (e.g., Promacta), or immunoglobulin (IVIg or anti-D) OR
  - B. Patient has an intolerance or hypersensitivity to a corticosteroid, another thrombopoietin receptor agonist (e.g., Promacta), or immunoglobulin (IVIg or anti-D) OR
  - C. Patient has an FDA labeled contraindication to a corticosteroid, another thrombopoietin receptor agonist (e.g., Promacta), or immunoglobulin (IVIg or anti-D) OR
  - D. Patient has had an insufficient response to a splenectomy AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of chronic immune (idiopathic) thrombocytopenia (ITP) AND ONE of the following:
  - A. Patient's platelet count is 50 x 10^9/L or greater OR
  - B. Patient's platelet count has increased sufficiently to avoid clinically significant bleeding AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

### Age Restriction:

**Prescriber Restrictions:** 

#### **Coverage Duration:**

Approval will be 6 months for initial, 12 months for renewal

Tavneos PA

### Drug Name(s)

Tavneos

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

Severe hepatic impairment (Child-Pugh C)

### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of severe active anti-neutrophil cytoplasmic autoantibody (ANCA)-associated vasculitis (granulomatosis with polyangiitis [GPA] and/or microscopic polyangiitis [MPA]) AND
- 2. Patient will continue standard therapy (e.g., corticosteroids, azathioprine, mycophenolate mofetil) in combination with the requested agent for the requested indication

## Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., rheumatologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Tegsedi PA

### Drug Name(s)

Tegsedi

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of polyneuropathy of hereditary transthyretin-mediated amyloidosis AND
- 2. The diagnosis has been confirmed by biopsy or genetic testing AND
- 3. Patient has clinical manifestations of polyneuropathy (e.g., neuropathic pain, altered sensation, numbness, tingling, impaired balance, motor disability) AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of polyneuropathy of hereditary transthyretin-mediated amyloidosis AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

#### Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., geneticist, neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Teriparatide PA

#### Drug Name(s)

Forteo

Teriparatide

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

#### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has ONE of the following:
  - A. Postmenopausal osteoporosis OR
  - B. Patient's sex is male with primary or hypogonadal osteoporosis OR
  - C. Osteoporosis with sustained systemic glucocorticoid therapy AND
- 2. Patient's diagnosis was confirmed by ONE of the following:
  - A. A fragility fracture in the hip or spine OR
  - B. A T-score of -2.5 or lower OR
  - C. A T-score of -1.0 to -2.5 AND ONE of the following:
    - i. A fragility fracture of the proximal humerus, pelvis, or distal forearm OR
    - ii. A FRAX 10-year probability for major osteoporotic fracture of 20% or greater OR
    - iii. A FRAX 10-year probability of hip fracture of 3% or greater AND
- 3. ONE of the following:
  - A. Patient is at a very high fracture risk as defined by ONE of the following:
    - i. Patient had a recent fracture (within the past 12 months) OR
    - ii. Patient had fractures while on FDA approved osteoporosis therapy OR
    - iii. Patient has had multiple fractures OR
    - iv. Patient had fractures while on drugs causing skeletal harm (e.g., long-term glucocorticoids) OR
    - v. Patient has a very low T-score (less than -3.0) OR
    - vi. Patient is at high risk for falls or has a history of injurious falls OR
    - vii. Patient has a very high fracture probability by FRAX (e.g., major osteoporosis fracture greater than 30%, hip fracture greater than 4.5%) or by other validated fracture risk algorithm OR
  - B. ONE of the following:
    - i. Patient has tried and had an inadequate response to a bisphosphonate OR
    - ii. Patient has an intolerance or hypersensitivity to a bisphosphonate OR
    - iii. Patient has an FDA labeled contraindication to a bisphosphonate AND
- 4. Patient will NOT be using the requested agent in combination with a bisphosphonate, denosumab (e.g., Prolia, Xgeva), romosozumab-aqqg, or another parathyroid hormone analog (e.g., abaloparatide) for the requested indication AND

Criteria continues: see Other Criteria

**Age Restriction:** 

#### **Prescriber Restrictions:**

### **Coverage Duration:**

No prior teriparatide and/or Tymlos use approve 2 years, Prior use - see Other Criteria Other Criteria:

- 5. The requested dose is within FDA labeled dosing for the requested indication AND 6. ONE of the following:
  - A. Patient has never received treatment with teriparatide or Tymlos (abaloparatide) OR
    - B. Patient has been previously treated with teriparatide or Tymlos (abaloparatide) AND ONE of the following:
      - i. The total cumulative duration of treatment with teriparatide and Tymlos (abaloparatide) has NOT exceeded 2 years OR
      - ii. Patient has received 2 years or more of treatment with teriparatide, or a combination of teriparatide and Tymlos (abaloparatide), and remains at or has returned to having a high risk for fracture

Prior teriparatide and/or Tymlos use approve remainder of 2 years of total cumulative therapy. Approve 1 year if patient has received 2 years or more teriparatide or a combination of teriparatide and Tymlos (abaloparatide)

Tetrabenazine PA

### Drug Name(s)

Tetrabenazine

Xenazine

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

#### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of chorea associated with Huntington's disease OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. Patient does NOT have a current diagnosis of depression OR
  - B. Patient has a current diagnosis of depression and is being treated for depression AND
- 3. ONE of the following:
  - A. Patient does NOT have a diagnosis of suicidal ideation and/or behavior OR
  - B. Patient has a diagnosis of suicidal ideation and/or behavior and must NOT be actively suicidal AND
- 4. Patient will NOT be using the requested agent in combination with a monoamine oxidase inhibitor (MAOI) AND
- 5. Patient will NOT be using the requested agent in combination with reserpine

#### Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 months

Tezspire PA

### Drug Name(s)

Tezspire

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of severe asthma AND
- 2. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LRTA, LAMA, theophylline) in combination with the requested agent AND
- 3. Patient will NOT be using the requested agent in combination with Xolair, an IL-5 inhibitor (Cinqair, Fasenra, Nucala), or Dupixent for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of severe asthma AND
- 3. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LRTA, LAMA, theophylline) in combination with the requested agent AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with Xolair, an IL-5 inhibitor (Cinqair, Fasenra, Nucala), or Dupixent for the requested indication

### Age Restriction:

Patient is 12 years of age or over

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, immunologist, pulmonologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# Coverage Duration:

Approval will be for 12 months

Tlando PA

### Drug Name(s)

Tlando

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient's sex is male with a diagnosis of primary or secondary (hypogonadotropic) hypogonadism AND
- 2. ONE of the following:
  - A. Patient is NOT currently receiving testosterone replacement therapy AND has ONE of the following pretreatment levels:
    - i. Total serum testosterone level that is below the testing laboratory's lower limit of the normal range or is less than 300 ng/dL OR
    - ii. Free serum testosterone level that is below the testing laboratory's lower limit of the normal range OR
  - B. Patient is currently receiving testosterone replacement therapy AND has ONE of the following current levels:
    - i. Total serum testosterone level that is within the testing laboratory's normal range OR below the testing laboratory's lower limit of the normal range OR is less than 300 ng/dL OR
    - ii. Free serum testosterone level is within the testing laboratory's normal range OR below the testing laboratory's normal range AND
- 3. ONE of the following:
  - A. Patient will NOT be using the requested agent in combination with another androgen or anabolic steroid OR
  - B. Prescriber has provided information in support of therapy with more than one agent

### Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 months

Tobi Podhaler PA

Drug Name(s)

Tobi Podhaler

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Tobramycin neb PA

Drug Name(s)

Kitabis Pak

Tobi

Tobramycin Neb

**Indications:** 

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

**Required Medical Information:** 

Pending CMS Review

Topical Diclofenac 3% Gel PA

Drug Name(s)

Diclofenac Sodium Gel 3%

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has a diagnosis of actinic keratosis (AK)

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 3 months

Topical Doxepin PA

# Drug Name(s)

Doxepin Hcl

Prudoxin

Zonalon

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - a. Patient has a diagnosis of moderate pruritus associated with atopic dermatitis AND ONE of the following:
    - i. Patient has tried and had an inadequate response to a topical corticosteroid (e.g., hydrocortisone, triamcinolone) OR
    - ii. Patient has an intolerance or hypersensitivity to a topical corticosteroid OR
    - iii. Patient has an FDA labeled contraindication to a topical corticosteroid OR
  - b. Patient has a diagnosis of moderate pruritus associated with lichen simplex chronicus AND ONE of the following:
    - i. Patient has tried and had an inadequate response to a topical corticosteroid (e.g., hydrocortisone, triamcinolone) OR
    - ii. Patient has an intolerance or hypersensitivity to a topical corticosteroid OR
    - iii. Patient has an FDA labeled contraindication to a topical corticosteroid

#### Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 8 days

Topical NSAID PA – Flector

# Drug Name(s)

Diclofenac Epolamine

Flector

#### Indications:

All Medically-Accepted Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - a. Patient has an FDA labeled indication for the requested agent OR
  - b. Patient has an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

### **Prescriber Restrictions:**

### **Coverage Duration:**

3 months for acute pain, 12 months for all other diagnoses

Topical NSAID PA – Licart

# Drug Name(s)

Licart

#### Indications:

All Medically-Accepted Indications.

### **Off-Label Uses:**

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - a. Patient has an FDA labeled indication for the requested agent OR
  - b. Patient has an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

# **Prescriber Restrictions:**

## **Coverage Duration:**

3 months for acute pain, 12 months for all other diagnoses

Topical NSAID PA - Pennsaid

### Drug Name(s)

Diclofenac Sodium (Pennsaid)

Pennsaid

#### Indications:

All Medically-Accepted Indications.

### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - a. Patient has an FDA labeled indication for the requested agent OR
  - b. Patient has an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

### **Prescriber Restrictions:**

### **Coverage Duration:**

3 months for acute pain, 12 months for all other diagnoses

Topical Retinoids PA – Adapalene

# Drug Name(s)

Adapalene

Adapalene Pump

Differin

### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

Requested agent will be used for cosmetic purposes

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - a. Patient has an FDA labeled indication for the requested agent OR
  - b. Patient has an indication that is supported in CMS approved compendia for the requested agent

### Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Topical Retinoids PA – Tazarotene

# Drug Name(s)

Arazlo

Fabior

Tazarotene

Tazorac

### Indications:

All Medically-Accepted Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

Requested agent will be used for cosmetic purposes

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - a. Patient has an FDA labeled indication for the requested agent OR
  - b. Patient has an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

**Prescriber Restrictions:** 

# **Coverage Duration:**

Approval will be for 12 months

Topical Retinoids PA – Tretinoin

# Drug Name(s)

Altreno

Atralin

Avita

Retin-A

Retin-A Micro

Retin-A Micro Pump

Tretinoin Cream, Gel

Tretinoin Microsphere

Tretinoin Microsphere Pump

### **Indications:**

All Medically-Accepted Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

Requested agent will be used for cosmetic purposes

### **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - a. Patient has an FDA labeled indication for the requested agent OR
  - b. Patient has an indication that is supported in CMS approved compendia for the requested agent

# Age Restriction:

### **Prescriber Restrictions:**

### **Coverage Duration:**

Approval will be for 12 months

Topical Retinoids PA – Trifarotene

# Drug Name(s)

Aklief

### Indications:

All Medically-Accepted Indications.

### **Off-Label Uses:**

### **Exclusion Criteria:**

Requested agent will be used for cosmetic purposes

# **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - a. Patient has an FDA labeled indication for the requested agent OR
  - b. Patient has an indication that is supported in CMS approved compendia for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Trelstar PA

## Drug Name(s)

Trelstar Mixject

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. BOTH of the following:
    - i. Patient is NOT currently being treated with the requested agent AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent AND
- 3. The requested dose is within FDA labeled dosing or supported in CMS approved compendia dosing for the requested indication

**Age Restriction:** 

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Trientine PA

### Drug Name(s)

Syprine

Trientine Hcl

#### Indications:

All FDA-Approved Indications.

# Off-Label Uses:

#### **Exclusion Criteria:**

# **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of Wilson's disease confirmed by ONE of the following:
  - A. Confirmation of genetic mutation of the ATP7B gene OR
  - B. Patient has TWO or more of the following:
    - i. Presence of hepatic abnormality (e.g., acute liver failure, cirrhosis, fatty liver)
    - ii. Presence of Kayser-Fleischer rings
    - iii. Serum ceruloplasmin level less than 20 mg/dL
    - iv. Basal urinary copper excretion greater than 40 mcg/24 hours or the testing laboratory's upper limit of normal
    - v. Hepatic parenchymal copper content greater than 40 mcg/g dry weight
    - vi. Presence of neurological symptoms (e.g., dystonia, hypertonia, rigidity with tremors, dysarthria, muscle spasms, dysphasia, polyneuropathy, dysautonomia) AND
- 2. ONE of the following:
  - A. Patient has tried and had an inadequate response to penicillamine OR
  - B. Patient has an intolerance or hypersensitivity to penicillamine OR
  - C. Patient has an FDA labeled contraindication to penicillamine

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of Wilson's disease AND
- 3. Patient has had clinical benefit with the requested agent as evidenced by ONE of the following:
  - A. Improvement and/or stabilization in hepatic abnormality OR
  - B. Reduction in Kayser-Fleischer rings OR
  - C. Improvement and/or stabilization in neurological symptoms (e.g., dystonia, hypertonia, rigidity with tremors, dysarthria, muscle spasms, dysphasia, polyneuropathy, dysautonomia) OR
  - D. Basal urinary copper excretion greater than 200 mcg/24 hours

## **Age Restriction:**

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, hepatologist, neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Approval will be for 12 months

Trikafta PA

## Drug Name(s)

Trikafta

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of cystic fibrosis AND
- 2. ONE of the following:
  - A. Patient has the presence of the F508del mutation in at least ONE allele (heterozygous OR homozygous) of the CFTR gene confirmed by genetic testing OR
  - B. Patient has ONE of the CFTR gene mutations or a mutation in the CFTR gene that is responsive based on in vitro data, as indicated in the FDA label, confirmed by genetic testing OR
  - C. Patient has another CFTR gene mutation(s) that is responsive to the requested agent, as indicated in the FDA label, confirmed by genetic testing AND
- 3. Patient will NOT be using the requested agent in combination with another CFTR modulator agent for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of cystic fibrosis AND
- 3. Patient has had improvement or stabilization with the requested agent [e.g., improvement in FEV1 from baseline, increase in weight/BMI, improvement from baseline Cystic Fibrosis Questionnaire-Revised (CFQ-R) Respiratory Domain score, improvements in respiratory symptoms (cough, sputum production, and difficulty breathing), and/or reduced number of pulmonary exacerbations] AND
- 4. Patient will NOT be using the requested agent in combination with another CFTR modulator agent for the requested indication

#### Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cystic fibrosis, pulmonologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Trudhesa PA

## Drug Name(s)

Trudhesa

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. The requested agent will be used for the treatment of acute migraine with or without aura AND
- 2. ONE of the following:
  - A. Patient has tried and had an inadequate response to TWO triptan agents with differing active ingredients (e.g., sumatriptan, rizatriptan) OR
  - B. Patient has an intolerance or hypersensitivity to TWO triptan agents with differing active ingredients OR
  - C. Patient has an FDA labeled contraindication to TWO triptan agents with differing active ingredients AND
- 3. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, acute CGRP)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. The requested agent will be used for the treatment of acute migraine with or without aura AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, acute CGRP)

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Trulicity PA

## Drug Name(s)

Trulicity

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

Requested agent will be used for weight loss alone

### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of type 2 diabetes mellitus AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 180 days OR
  - C. ALL of the following:
    - i. ONE of the following:
      - 1. Patient's medication history includes use of a non glucagon-like peptide-1 (GLP-1) oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) within the past 90 days OR
      - 2. Patient had an ineffective treatment response to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 3. Patient has an intolerance or hypersensitivity to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 4. Patient has an FDA labeled contraindication to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 5. BOTH of the following:
        - a. Patient has a diagnosis of established cardiovascular disease [e.g., myocardial infarction, stroke, any revascularization procedure, transient ischemic attack, unstable angina, amputation, symptomatic or asymptomatic coronary artery disease] AND
        - b. The requested agent will be used to reduce the risk of major adverse cardiovascular events (cardiovascular death, non-fatal myocardial infarction, or non-fatal stroke) AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent AND
    - iii. Patient will NOT be using the requested agent in combination with another GLP-1 agonist agent, or an agent containing a GLP-1 agonist AND
    - iv. Patient will NOT be using the requested agent in combination with an agent containing a dipeptidyl peptidase-4 (DPP-4) inhibitor

**Age Restriction:** 

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Other Criteria:

Tymlos PA

### Drug Name(s)

**Tymlos** 

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient (pt) has ONE of the following:
  - A. Postmenopausal osteoporosis OR
  - B. Pt's sex is male with osteoporosis AND
- 2. BOTH of the following:
  - A. Pt's diagnosis was confirmed by ONE of the following:
    - i. A fragility fracture in the hip or spine OR
    - ii. A T-score of -2.5 or lower OR
    - iii. A T-score of -1.0 to -2.5 AND ONE of the following:
      - a. A fragility fracture of proximal humerus, pelvis, or distal forearm OR
      - b. A FRAX 10-year probability for major osteoporotic fracture of 20% or greater OR
      - c. A FRAX 10-year probability of hip fracture of 3% or greater AND
  - B. ONE of the following:
    - i. Pt is at a very high fracture risk as defined by ONE of the following:
      - a. Pt had a recent fracture (within the past 12 months) OR
      - b. Pt had fractures while on FDA approved osteoporosis therapy OR
      - c. Pt has had multiple fractures OR
      - d. Pt had fractures while on drugs causing skeletal harm (e.g., long-term glucocorticoids) OR
      - e. Pt has a very low T-score (less than -3.0) OR
      - f. Pt is at high risk for falls or has a history of injurious falls OR
      - g. Pt has a very high fracture probability by FRAX (e.g., major osteoporosis fracture greater than 30%, hip fracture greater than 4.5%) or by other validated fracture risk algorithm OR
    - ii. ONE of the following:
      - a. Pt has tried and had an inadequate response to a bisphosphonate OR
      - b. Pt has an intolerance or hypersensitivity to a bisphosphonate OR
      - c. Pt has an FDA labeled contraindication to a bisphosphonate AND
- 3. Pt will NOT be using the requested agent in combination with a bisphosphonate, denosumab (e.g., Prolia, Xgeva), romosozumab-aqqg, or another parathyroid hormone analog (e.g., teriparatide) for the requested indication AND
- 4. The requested dose is within FDA labeled dosing for the requested indication AND
- 5. The total cumulative duration of treatment with teriparatide and Tymlos (abaloparatide) has not exceeded 2 years

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

No prior Tymlos and/or teriparatide use approve 2 years, Prior use - see Other Criteria

Other Criteria:

Prior Tymlos and/or teriparatide use approve remainder of 2 years of total cumulative therapy

Tyrvaya PA

# Drug Name(s)

Tyrvaya

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Ubrelvy PA

## Drug Name(s)

Ubrelvy

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of migraine AND
- 2. The requested agent is being used for the treatment of acute migraine with or without aura AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to a triptan (e.g., sumatriptan, rizatriptan) agent OR
  - B. Patient has an intolerance, or hypersensitivity to a triptan OR
  - C. Patient has an FDA labeled contraindication to a triptan AND
- 4. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, ergotamine, acute CGRP)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of migraine AND
- 3. The requested agent is being used for the treatment of acute migraine with or without aura AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, ergotamine, acute CGRP)

### Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 months

Urea Cycle Disorders PA - Olpruva

## Drug Name(s)

Olpruva

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of urea cycle disorders (UCDs), involving deficiencies of carbamylphosphate synthetase (CPS), ornithine transcarbamylase (OTC), or argininosuccinic acid synthetase (AS) AND
- 2. The requested agent will be used as chronic management of UCDs AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

## **Age Restriction:**

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., geneticist, metabolic disorders) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Urea Cycle Disorders PA – Pheburane

## Drug Name(s)

Pheburane

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of urea cycle disorders (UCDs), involving deficiencies of carbamylphosphate synthetase (CPS), ornithine transcarbamylase (OTC), or argininosuccinic acid synthetase (AS) AND
- 2. The requested agent will be used as chronic management of UCDs AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

## Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., geneticist, metabolic disorders) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Urea Cycle Disorders PA – Ravicti

### Drug Name(s)

Ravicti

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of urea cycle disorder and the requested agent will be used for chronic management AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

### **Age Restriction:**

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., geneticist, metabolic disorders) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Urea Cycle Disorders PA - Sodium Phenylbutyrate

# Drug Name(s)

Buphenyl

Sodium Phenylbutyrate

#### **Indications:**

All FDA-Approved Indications.

### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of ONE of the following:
  - a. Urea cycle disorder with neonatal-onset involving deficiencies of carbamylphosphate synthetase, ornithine transcarbamylase, or argininosuccinic acid synthetase OR
  - b. Urea cycle disorder with late-onset and history of hyperammonemic encephalopathy involving deficiencies of carbamylphosphate synthetase, ornithine transcarbamylase, or argininosuccinic acid synthetase AND
- 2. The requested dose is within FDA labeled dosing for the requested indication

# Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., geneticist, metabolic disorders) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Valchlor PA

### Drug Name(s)

Valchlor

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. BOTH of the following:
    - i. ONE of the following:
      - a. BOTH of the following:
        - 1. Patient has a diagnosis of Stage IA or IB mycosis fungoides-type cutaneous T-cell lymphoma AND
        - 2. Patient's medication history indicates use of at least ONE prior skindirected therapy (e.g., topical corticosteroid) OR
      - b. Patient has an indication that is supported in CMS approved compendia for the requested agent AND
    - ii. Prescriber is a specialist in the area of the patient's diagnosis (e.g., dermatologist, oncologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has an FDA labeled indication or an indication that is supported in CMS approved compendia for the requested agent AND
- 3. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. BOTH of the following:
    - i. Patient has had clinical benefit with the requested agent AND
    - ii. Prescriber is a specialist in the area of the patient's diagnosis (e.g., dermatologist, oncologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### Age Restriction:

Prescriber Restrictions:
Coverage Duration:
Approval will be for 12 months
Other Criteria:

Veozah PA

# Drug Name(s)

Veozah

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

V-Go PA

### Drug Name(s)

V-Go

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of diabetes mellitus AND
- 2. Patient is on an insulin regimen of 3 or more injections per day AND
- 3. ONE of the following:
  - A. Patient is testing glucose levels 4 or more times per day OR
  - B. Patient is using a continuous glucose monitor (CGM)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of diabetes mellitus AND
- 3. Patient has had clinical benefit with the requested agent (e.g., stable or improved glycemic control)

### Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 12 months

Viberzi PA

# Drug Name(s)

Viberzi

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Criteria for approval require the following:

1. Patient has a diagnosis of irritable bowel syndrome with diarrhea (IBS-D)

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Victoza PA

## Drug Name(s)

Liraglutide

Victoza

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

Requested agent will be used for weight loss alone

### **Required Medical Information:**

Criteria for approval require BOTH of the following:

- 1. Patient has a diagnosis of type 2 diabetes mellitus AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 180 days OR
  - C. ALL of the following:
    - i. ONE of the following:
      - 1. Patient's medication history includes use of a non glucagon-like peptide-1 (GLP-1) oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) within the past 90 days OR
      - 2. Patient had an ineffective treatment response to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 3. Patient has an intolerance or hypersensitivity to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR
      - 4. Patient has an FDA labeled contraindication to a non GLP-1 oral diabetes medication (e.g., metformin, an agent containing metformin, glipizide) OR 5. BOTH of the following:
        - a. Patient has a diagnosis of established cardiovascular disease [e.g., myocardial infarction, stroke, any revascularization procedure, transient ischemic attack, unstable angina, amputation, symptomatic or asymptomatic coronary artery disease] AND
        - b. The requested agent will be used to reduce the risk of major adverse cardiovascular events (cardiovascular death, non-fatal myocardial infarction, or non-fatal stroke) AND
    - ii. Patient does NOT have any FDA labeled contraindications to the requested agent AND iii. Patient will NOT be using the requested agent in combination with another GLP-1
    - agonist agent, or an agent containing a GLP-1 agonist AND
    - iv. Patient will NOT be using the requested agent in combination with an agent containing a dipeptidyl peptidase-4 (DPP-4) inhibitor

### Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Vijoice PA

# Drug Name(s)

Vijoice

### **Indications:**

All FDA-Approved Indications.

#### **Off-Label Uses:**

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of PIK3CA-Related Overgrowth Spectrum (PROS) confirmed by genetic testing indicating a mutation in the PIK3CA gene AND
- 2. Patient has severe manifestations of PROS AND
- 3. Patient requires systemic therapy

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of PIK3CA-Related Overgrowth Spectrum (PROS) AND
- 3. Patient has severe manifestations of PROS AND
- 4. Patient requires systemic therapy AND
- 5. Patient has had clinical benefit with the requested agent

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

### **Prescriber Restrictions:**

# **Coverage Duration:**

Approval will be for 12 months

Vivjoa PA

## Drug Name(s)

Vivjoa

### **Indications:**

All FDA-Approved Indications.

#### **Off-Label Uses:**

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of vulvovaginal candidiasis AND
- 2. The requested agent will be used to reduce the incidence of recurrent vulvovaginal candidiasis AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to fluconazole OR
  - B. Patient has an intolerance or hypersensitivity to fluconazole OR
  - C. Patient has an FDA labeled contraindication to fluconazole OR
  - D. Patient will be using fluconazole as part of the fluconazole/Vivjoa dosage regimen

### Age Restriction:

**Prescriber Restrictions:** 

### **Coverage Duration:**

Approval will be for 4 months

Voriconazole PA

#### Drug Name(s)

Vfend

Vfend Iv

Voriconazole

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of invasive Aspergillus OR
  - B. Patient has a serious infection caused by Scedosporium apiospermum or Fusarium species OR
  - C. Patient has a diagnosis of esophageal candidiasis or candidemia in nonneutropenic patient AND ONE of the following:
    - i. Patient has tried and had an inadequate response to fluconazole or an alternative antifungal agent OR
    - ii. Patient has an intolerance or hypersensitivity to fluconazole or an alternative antifungal agent OR
    - iii. Patient has an FDA labeled contraindication to fluconazole or an alternative antifungal agent OR
  - D. Patient has a diagnosis of blastomycosis AND ONE of the following:
    - i. Patient has tried and had an inadequate response to itraconazole OR
    - ii. Patient has an intolerance or hypersensitivity to itraconazole OR
    - iii. Patient has an FDA labeled contraindication to itraconazole OR
  - E. The requested agent is being prescribed for prophylaxis of invasive Aspergillus or Candida AND patient is severely immunocompromised, such as a hematopoietic stem cell transplant [HSCT] recipient, or hematologic malignancy with prolonged neutropenia from chemotherapy, or is a high-risk solid organ (lung, heart-lung, liver, pancreas, small bowel) transplant patient, or long term use of high dose corticosteroids (greater than 1 mg/kg/day of prednisone or equivalent) OR
  - F. Patient has another indication that is supported in CMS approved compendia for the requested agent

### **Age Restriction:**

## **Prescriber Restrictions:**

### **Coverage Duration:**

One month for esophageal candidiasis, 6 months for all other indications

#### Other Criteria:

Criteria for renewal approval require BOTH of the following:

1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND

### 2. ONE of the following:

A. Patient has a diagnosis of invasive Aspergillus, a serious infection caused by Scedosporium apiospermum or Fusarium species, esophageal candidiasis, candidemia in nonneutropenic patient, or blastomycosis and patient has continued indicators of active disease (e.g., continued radiologic findings, positive cultures, positive serum galactomannan assay for Aspergillus) OR B. The requested agent is being prescribed for prophylaxis of invasive Aspergillus or Candida and patient continues to be severely immunocompromised, such as a hematopoietic stem cell transplant [HSCT] recipient, or hematologic malignancy with prolonged neutropenia from chemotherapy, or is a high-risk solid organ (lung, heart-lung, liver, pancreas, small bowel) transplant patient, or long term use of high dose corticosteroids (greater than 1 mg/kg/day of prednisone or equivalent) OR

### C. BOTH of the following:

- i. Patient has another indication that is supported in CMS approved compendia for the requested agent  $\ensuremath{\mathsf{AND}}$
- ii. Patient has had clinical benefit with the requested agent

Vosevi PA

### Drug Name(s)

Vosevi

#### Indications:

All Medically-Accepted Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of hepatitis C confirmed by serological markers AND
- 2. Prescriber has screened the patient for current or prior hepatitis B viral (HBV) infection and if positive, will monitor the patient for HBV flare-up or reactivation during and after treatment with the requested agent AND
- 3. The requested agent will be used in a treatment regimen and length of therapy that is supported in FDA approved labeling or AASLD/IDSA guidelines for the patient's diagnosis and genotype AND
- 4. The requested dose is within FDA labeled dosing or supported in AASLD/IDSA guideline dosing for the requested indication AND
- 5. If genotype 1, the patient's subtype has been identified and provided

### **Age Restriction:**

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, hepatologist or infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

# **Coverage Duration:**

Duration of therapy: Based on FDA approved labeling or AASLD/IDSA guideline supported

Vowst PA

### Drug Name(s)

Vowst

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. The requested agent will be used to prevent the recurrence of Clostridioides difficile infection (CDI) AND
- 2. Patient has had a confirmed diagnosis of recurrent CDI as defined by greater than or equal to 3 episodes of CDI in a 12 month period AND
- 3. Patient has completed a standard of care antibiotic regimen (e.g., vancomycin, fidaxomicin) for recurrent CDI at least 2 to 4 days before initiating treatment with the requested agent AND
- 4. Patient will NOT be using the requested agent in combination with any antibiotic regimen for any indication

### Age Restriction:

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., infectious disease, gastroenterologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis **Coverage Duration:** 

Approval will be for 12 months

Voxzogo PA

## Drug Name(s)

Voxzogo

### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of achondroplasia AND
- 2. The requested agent will be used to increase linear growth AND
- 3. Patient has open epiphyses

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of achondroplasia AND
- 3. The requested agent will be used to increase linear growth AND
- 4. Patient has open epiphyses AND
- 5. Patient has had clinical benefit with the requested agent

## Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., endocrinologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Voydeya PA

# Drug Name(s)

Voydeya

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of paroxysmal nocturnal hemoglobinuria (PNH) AND
- 2. The requested agent will be used for the treatment of extravascular hemolysis (EVH) AND
- 3. The requested agent will be used in combination with Soliris (eculizumab) or Ultomiris (ravulizumab-cwvz)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of paroxysmal nocturnal hemoglobinuria (PNH) AND
- 3. The requested agent will be used for the treatment of extravascular hemolysis (EVH) AND
- 4. The requested agent will be used in combination with Soliris (eculizumab) or Ultomiris (ravulizumab-cwvz) AND
- 5. Patient has had clinical benefit with the requested agent [e.g., decreased requirement for packed red blood cell transfusions, stabilization/improvement of hemoglobin]

### **Age Restriction:**

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., hematologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Vtama PA

## Drug Name(s)

Vtama

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of plaque psoriasis AND
- 2. ONE of the following:
  - a. Patient has tried and failed a topical corticosteroid (e.g., triamcinolone) OR
  - b. Patient has an intolerance or hypersensitivity to a topical steroid OR
  - c. Patient has an FDA labeled contraindication to a topical steroid AND
- 3. ONE of the following:
  - a. Patient has tried and failed a topical vitamin D analog (e.g., calcipotriene) OR tazarotene OR
  - b. Patient has an intolerance or hypersensitivity to a topical vitamin D analog OR tazarotene OR
  - c. Patient has an FDA labeled contraindication to a topical vitamin D analog OR tazarotene

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of plaque psoriasis AND
- 3. Patient has had clinical benefit with the requested agent

## Age Restriction:

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., dermatologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Vyndamax PA

# Drug Name(s)

Vyndamax

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of cardiomyopathy of wild type or hereditary transthyretin-mediated amyloidosis (ATTR-CM) AND
- 2. The diagnosis has been confirmed by testing [e.g., stannous pyrophosphate (PYP) scanning, monoclonal antibody studies, biopsy, scintigraphy, genetic testing] AND
- 3. The requested agent will be used to reduce cardiovascular mortality and cardiovascular-related hospitalization AND
- 4. Patient will NOT be using the requested agent in combination with another tafamidis agent for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of cardiomyopathy of wild type or hereditary transthyretin-mediated amyloidosis (ATTR-CM) AND
- 3. The requested agent will be used to reduce cardiovascular mortality and cardiovascular-related hospitalization AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with another tafamidis agent for the requested indication

### Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cardiologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Vyndagel PA

## Drug Name(s)

Vyndagel

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of cardiomyopathy of wild type or hereditary transthyretin-mediated amyloidosis (ATTR-CM) AND
- 2. The diagnosis has been confirmed by testing [e.g., stannous pyrophosphate (PYP) scanning, monoclonal antibody studies, biopsy, scintigraphy, genetic testing] AND
- 3. The requested agent will be used to reduce cardiovascular mortality and cardiovascular-related hospitalization AND
- 4. Patient will NOT be using the requested agent in combination with another tafamidis agent for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of cardiomyopathy of wild type or hereditary transthyretin-mediated amyloidosis (ATTR-CM) AND
- 3. The requested agent will be used to reduce cardiovascular mortality and cardiovascular-related hospitalization AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with another tafamidis agent for the requested indication

### Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cardiologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Wainua PA

## Drug Name(s)

Wainua

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

### **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

### **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of polyneuropathy of hereditary transthyretin-mediated amyloidosis AND
- 2. The diagnosis has been confirmed by biopsy or genetic testing AND
- 3. Patient has clinical manifestations of polyneuropathy (e.g., neuropathic pain, altered sensation, numbness, tingling, impaired balance, motor disability) AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of polyneuropathy of hereditary transthyretin-mediated amyloidosis AND
- 3. Patient has had clinical benefit with the requested agent AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

### Age Restriction:

### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., geneticist, neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

### **Coverage Duration:**

Approval will be for 12 months

Wakix PA

Drug Name(s)

Wakix

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Pending CMS Review

Wegovy PA

# Drug Name(s)

Wegovy

# Indications:

All FDA-Approved Indications.

## Off-Label Uses:

## **Exclusion Criteria:**

Requested agent will be used for weight loss alone AND FDA labeled contraindications to the requested agent

# **Required Medical Information:**

Pending CMS Review

Winlevi PA

## Drug Name(s)

Winlevi

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

Requested agent will be used for cosmetic purposes

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has a diagnosis of acne vulgaris

Age Restriction:

Patient is within the FDA labeled age for the requested agent

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Xdemvy PA

Drug Name(s)

Xdemvy

**Indications:** 

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

**Required Medical Information:** 

Criteria for approval require the following:

1. Patient has an FDA labeled indication for the requested agent

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 6 weeks

Xembify PA

Drug Name(s)

Xembify

**Indications:** 

All Medically-Accepted Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require ONE of the following:

- 1. Patient has ONE of the following diagnoses:
  - A. Primary immunodeficiency [e.g., congenital agammaglobulinemia, common variable immunodeficiency (CVID), severe combined immunodeficiency, Wiskott-Aldrich Syndrome, X-linked agammaglobulinemia (XLA), humoral immunodeficiency, IgG subclass deficiency with or without IgA deficiency] OR
  - B. Multiple sclerosis (MS) AND BOTH of the following:
    - i. Patient has a diagnosis of relapsing remitting MS (RRMS) AND
    - ii. Patient has had an insufficient response, documented failure, or FDA labeled contraindication to TWO MS agents (e.g., Avonex, Betaseron, Copaxone, dimethyl fumarate, fingolimod, glatiramer, Glatopa, Kesimpta, Plegridy, teriflunomide, Vumerity) OR
- 2. ONE of the following:
  - A. Patient has another FDA labeled indication for the requested agent OR
  - B. Patient has an indication that is supported in CMS approved compendia for the requested agent

Drug is also subject to Part B versus Part D review.

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Xermelo PA

## Drug Name(s)

Xermelo

Indications:

All FDA-Approved Indications.

Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of carcinoid syndrome diarrhea AND
- 2. Patient has tried and had an inadequate response to treatment with a somatostatin analog (e.g., Sandostatin [octreotide], Sandostatin LAR [octreotide], Somatuline Depot [lanreotide]) AND
- 3. The requested agent will be used in combination with a somatostatin analog (e.g., Sandostatin [octreotide], Sandostatin LAR [octreotide], Somatuline Depot [lanreotide])

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of carcinoid syndrome diarrhea AND
- 3. Patient has had clinical benefit with the requested agent (e.g., reduction in the average number of daily bowel movements) AND
- 4. The requested agent will be used in combination with a somatostatin analog (e.g., Sandostatin [octreotide], Sandostatin LAR [octreotide], Somatuline Depot [lanreotide])

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Xgeva PA

## Drug Name(s)

Xgeva

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of multiple myeloma AND BOTH of the following:
    - i. The requested agent will be used for the prevention of skeletal-related events AND
    - ii. ONE of the following:
      - 1. Patient has a pretreatment or current calcium level that is NOT below the limits of the testing laboratory's normal range OR
      - 2. Patient has a pretreatment or current calcium level that is below the limits of the testing laboratory's normal range AND it will be corrected prior to use of the requested agent OR
      - 3. Prescriber has indicated that the patient is not at risk for hypocalcemia (not including risk associated with the requested agent) OR
  - B. Patient has a diagnosis of prostate cancer AND ALL of the following:
    - i. The requested agent will be used for the prevention of skeletal-related events AND
    - ii. Patient has bone metastases AND
    - iii. ONE of the following:
      - 1. Patient has a pretreatment or current calcium level that is NOT below the limits of the testing laboratory's normal range OR
      - 2. Patient has a pretreatment or current calcium level that is below the limits of the testing laboratory's normal range AND it will be corrected prior to use of the requested agent OR
      - 3. Prescriber has indicated that the patient is not at risk for hypocalcemia (not including risk associated with the requested agent) OR

Criteria continues: see Other Criteria

Age Restriction:

**Prescriber Restrictions:** 

## **Coverage Duration:**

Approval will be for 12 months

- C. Patient has a solid tumor cancer diagnosis (e.g., thyroid, non-small cell lung, kidney cancer, or breast cancer) AND ALL of the following:
  - i. The requested agent will be used for the prevention of skeletal-related events AND
  - ii. Patient has bone metastases AND

- iii. ONE of the following:
  - 1. Patient has a pretreatment or current calcium level that is NOT below the limits of the testing laboratory's normal range OR
  - 2. Patient has a pretreatment or current calcium level that is below the limits of the testing laboratory's normal range AND it will be corrected prior to use of the requested agent OR
  - 3. Prescriber has indicated that the patient is not at risk for hypocalcemia (not including risk associated with the requested agent) OR
- D. Patient has a diagnosis of giant cell tumor of bone AND ONE of the following:
  - i. Patient has a pretreatment or current calcium level that is NOT below the limits of the testing laboratory's normal range OR
  - ii. Patient has a pretreatment or current calcium level that is below the limits of the testing laboratory's normal range AND it will be corrected prior to use of the requested agent OR
  - iii. Prescriber has indicated that the patient is not at risk for hypocalcemia (not including risk associated with the requested agent) OR
- E. Patient has a diagnosis of hypercalcemia of malignancy AND
- 2. Patient will NOT be using the requested agent in combination with Prolia (denosumab) AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Xifaxan PA

## Drug Name(s)

Xifaxan

## **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require the following:

- 1. Patient has ONE of the following:
  - a. A diagnosis of irritable bowel syndrome with diarrhea (IBS-D) OR
  - b. A diagnosis of hepatic encephalopathy [reduction in risk of overt hepatic encephalopathy (HE) recurrence] OR
  - c. BOTH of the following:
    - i. A diagnosis of traveler's diarrhea (TD) AND
    - ii. The traveler's diarrhea is caused by noninvasive strains of Escherichia coli

# Age Restriction:

For diagnosis of traveler's diarrhea (TD), patient is within the FDA labeled age for the requested agent

## **Prescriber Restrictions:**

## **Coverage Duration:**

Approval will be for 12 months

Xolair PA

## Drug Name(s)

Xolair

#### Indications:

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of moderate to severe persistent asthma AND ALL of the following:
    - i. ONE of the following:
      - a. Patient is 6 to less than 12 years of age AND BOTH of the following:
        - I. Patient's pretreatment IgE level is 30 IU/mL to 1300 IU/mL AND
        - II. Patient's weight is 20 kg to 150 kg OR
      - b. Patient is 12 years of age or over AND BOTH of the following:
        - I. Patient's pretreatment IgE level is 30 IU/mL to 700 IU/mL AND
        - II. Patient's weight is 30 kg to 150 kg AND
    - ii. Allergic asthma has been confirmed by a positive skin test or in vitro reactivity test to a perennial aeroallergen AND
    - iii. ONE of the following:
      - a. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LRTA, LAMA, theophylline) in combination with the requested agent OR
      - b. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to an asthma control therapy OR
  - B. Patient has a diagnosis of chronic idiopathic urticaria AND BOTH of the following:
    - i. Patient has had over 6 weeks of hives and itching AND
    - ii. ONE of the following:
      - a. Patient has tried and had an inadequate response to maximum tolerable H1 antihistamine therapy OR
      - b. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to H1 antihistamine therapy OR
  - C. Patient has a diagnosis of nasal polyps AND BOTH of the following:
    - i. ONE of the following:
      - a. Patient has tried and had an inadequate response to an intranasal corticosteroid OR
      - b. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to an intranasal corticosteroid AND
    - ii. ONE of the following:
      - a. The requested agent will be used in combination with an intranasal corticosteroid OR

b. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to an intranasal corticosteroid OR

Initial criteria continues: see Other Criteria

### Age Restriction:

For diagnosis of moderate to severe persistent asthma, patient is 6 years of age or over. For diagnosis of chronic idiopathic urticaria, patient is 12 years of age or over. For diagnosis of nasal polyps, patient is 18 years of age or over. For diagnosis of IgE-mediated food allergy, patient is 1 year of age or over.

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., allergist, immunologist, otolaryngologist, pulmonologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be 6 months for initial, 12 months for renewal

## Other Criteria:

- D. Patient has a diagnosis of IgE-mediated food allergy AND ALL of the following:
  - i. Patient is using the requested agent for the reduction of allergic reactions (Type I), including anaphylaxis, that may occur with accidental exposure to one or more foods AND
  - ii. IgE-mediated food allergy has been confirmed by an allergy diagnostic test (e.g., skin prick test, serum specific IgE test, oral food challenge) AND
  - iii. Patient will avoid known food allergens while treated with the requested agent AND
  - iv. The requested agent will NOT be used for the emergency treatment of allergic reactions, including anaphylaxis AND
- 2. Patient will NOT be using the requested agent in combination with Dupixent or an injectable Interleukin 5 (IL-5) inhibitor (e.g., Cinqair, Fasenra, Nucala) for the requested indication AND
- 3. The requested dose is within FDA labeled dosing for the requested indication

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. ONE of the following:
  - A. Patient has a diagnosis of moderate to severe persistent asthma AND BOTH of the following:
    - i. Patient has had clinical benefit with the requested agent AND
    - ii. ONE of the following:
      - a. Patient is currently being treated with AND will continue asthma control therapy (e.g., ICS, ICS/LABA, LRTA, LAMA, theophylline) in combination with the requested agent OR
      - b. Patient has an intolerance, FDA labeled contraindication, or hypersensitivity to an asthma control therapy OR
  - B. Patient has a diagnosis of chronic idiopathic urticaria AND the following:
    - a. Patient has had clinical benefit with the requested agent OR
  - C. Patient has a diagnosis of nasal polyps AND the following:
    - a. Patient has had clinical benefit with the requested agent OR

- D. Patient has a diagnosis of IgE-mediated food allergy AND ALL of the following:
  - a. Patient is using the requested agent for the reduction of allergic reactions (Type I), including anaphylaxis, that may occur with accidental exposure to one or more foods AND
  - b. Patient has had clinical benefit with the requested agent AND
  - c. Patient will avoid known food allergens while treated with the requested agent AND
  - d. The requested agent will NOT be used for the emergency treatment of allergic reactions, including anaphylaxis AND
- 3. Patient will NOT be using the requested agent in combination with Dupixent or an injectable interleukin 5 (IL-5) inhibitor (e.g., Cinqair, Fasenra, Nucala) for the requested indication AND
- 4. The requested dose is within FDA labeled dosing for the requested indication

Xolremdi PA

Drug Name(s)

Xolremdi

**Indications:** 

All FDA-Approved Indications.

**Off-Label Uses:** 

**Exclusion Criteria:** 

FDA labeled contraindications to the requested agent

**Required Medical Information:** 

Pending CMS Review

Xywav PA

## Drug Name(s)

Xywav

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require the following:

- 1. ONE of the following:
  - A. Patient has a diagnosis of narcolepsy with cataplexy OR
  - B. Patient has a diagnosis of narcolepsy with excessive daytime sleepiness AND BOTH of the following:
    - i. ONE of the following:
      - a. Patient is under 18 years of age OR
      - b. ONE of the following:
        - 1. Patient has tried and had an inadequate response to modafinil or armodafinil OR
        - 2. Patient has an intolerance or hypersensitivity to modafinil or armodafinil OR
        - 3. Patient has an FDA labeled contraindication to modafinil or armodafinil AND
    - ii. ONE of the following:
      - a. Patient has tried and had an inadequate response to ONE standard stimulant agent (e.g., methylphenidate) OR
      - b. Patient has an intolerance or hypersensitivity to ONE standard stimulant agent (e.g., methylphenidate) OR
      - c. Patient has an FDA labeled contraindication to ONE standard stimulant agent (e.g., methylphenidate) OR
  - C. Patient has a diagnosis of idiopathic hypersomnia OR
  - D. Patient has another indication that is supported in CMS approved compendia for the requested agent

## Age Restriction:

For diagnosis of narcolepsy with cataplexy, patient is 7 years of age or over. For diagnosis of narcolepsy with excessive daytime sleepiness, patient is 7 years of age or over. For diagnosis of idiopathic hypersomnia, patient is 18 years of age or over.

## **Prescriber Restrictions:**

## **Coverage Duration:**

Approval will be for 12 months

Zavzpret PA

## Drug Name(s)

Zavzpret

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of migraine AND
- 2. The requested agent is being used for the treatment of acute migraine with or without aura AND
- 3. ONE of the following:
  - A. Patient has tried and had an inadequate response to a triptan (e.g., sumatriptan, rizatriptan) agent OR
  - B. Patient has an intolerance, or hypersensitivity to a triptan OR
  - C. Patient has an FDA labeled contraindication to a triptan AND
- 4. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, ergotamine, acute CGRP)

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of migraine AND
- 3. The requested agent is being used for the treatment of acute migraine with or without aura AND
- 4. Patient has had clinical benefit with the requested agent AND
- 5. Patient will NOT be using the requested agent in combination with another acute migraine agent (e.g., triptan, 5HT-1F, ergotamine, acute CGRP)

Age Restriction:

**Prescriber Restrictions:** 

**Coverage Duration:** 

Approval will be for 12 months

Zepatier PA

## Drug Name(s)

Zepatier

#### **Indications:**

All Medically-Accepted Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for approval require ALL of the following:

- 1. Patient has a diagnosis of hepatitis C confirmed by serological markers AND
- 2. Prescriber has screened the patient for current or prior hepatitis B viral (HBV) infection and if positive, will monitor the patient for HBV flare-up or reactivation during and after treatment with the requested agent AND
- 3. The requested agent will be used in a treatment regimen and length of therapy that is supported in FDA approved labeling or AASLD/IDSA guidelines for the patient's diagnosis and genotype AND
- 4. The requested dose is within FDA labeled dosing or supported in AASLD/IDSA guideline dosing for the requested indication AND
- 5. If genotype 1, the patient's subtype has been identified and provided AND
- 6. If genotype 1a, the prescriber has tested the patient for the presence of virus with NS5A resistance-associated polymorphisms AND
- 7. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 90 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent within the past 90 days OR
  - C. Patient has an FDA labeled contraindication or hypersensitivity to the preferred agent: Mavyret for supported genotypes OR
  - D. Prescriber has provided information based on FDA approved labeling or AASLD/IDSA guidelines supporting the use of the non-preferred agent for the patient's diagnosis and genotype over the preferred agent: Mavyret for supported genotypes

## **Age Restriction:**

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., gastroenterologist, hepatologist or infectious disease) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Duration of therapy: Based on FDA approved labeling or AASLD/IDSA guideline supported **Other Criteria**:

Zilbrysq PA

## Drug Name(s)

Zilbrysq

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require the following:

- 1. Patient has BOTH of the following:
  - A. Diagnosis of generalized myasthenia gravis (gMG) AND
  - B. A positive serological test for anti-AChR antibodies

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of generalized myasthenia gravis (gMG) AND
- 3. Patient has had clinical benefit with the requested agent [e.g., stabilization/improvement of Myasthenia Gravis-Activities of Daily Living score (MG-ADL) or Quantitative MG score (QMG)]

## **Age Restriction:**

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Zokinvy PA

## Drug Name(s)

Zokinvy

#### **Indications:**

All FDA-Approved Indications.

## **Off-Label Uses:**

## **Exclusion Criteria:**

FDA labeled contraindications to the requested agent

## **Required Medical Information:**

Criteria for initial approval require the following:

- 1. ONE of the following:
  - A. BOTH of the following:
    - i. Patient has a diagnosis of Hutchinson-Gilford progeria syndrome (HGPS) AND
    - ii. Genetic testing has confirmed a pathogenic variant in the LMNA gene that results in production of progerin OR
  - B. Patient has a diagnosis of processing-deficient progeroid laminopathy AND ONE of the following:
    - i. Genetic testing has confirmed heterozygous LMNA mutation with progerin-like protein accumulation OR
    - ii. Genetic testing has confirmed homozygous or compound heterozygous ZMPSTE24 mutations

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of ONE of the following:
  - A. Hutchinson-Gilford progeria syndrome (HGPS) OR
  - B. Processing-deficient progeroid laminopathies with either: heterozygous LMNA mutation with progerin-like protein accumulation OR homozygous or compound heterozygous ZMPSTE24 mutations AND
- 3. Patient has had clinical benefit with the requested agent

## **Age Restriction:**

Patient is within the FDA labeled age for the requested agent

#### **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., cardiologist, geneticist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Zoryve Cream PA

## Drug Name(s)

Zoryve Cream

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for initial approval require ALL of the following:

- 1. Patient has a diagnosis of plaque psoriasis AND
- 2. ONE of the following:
  - A. Patient has tried and failed a topical corticosteroid (e.g., triamcinolone) OR
  - B. Patient has an intolerance or hypersensitivity to a topical corticosteroid OR
  - C. Patient has an FDA labeled contraindication to a topical corticosteroid AND
- 3. ONE of the following:
  - A. Patient has tried and failed a topical vitamin D analog (e.g., calcipotriene) OR tazarotene OR
  - B. Patient has an intolerance or hypersensitivity to a topical vitamin D analog OR tazarotene OR
  - C. Patient has an FDA labeled contraindication to a topical vitamin D analog OR tazarotene

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of plaque psoriasis AND
- 3. Patient has had clinical benefit with the requested agent

## **Age Restriction:**

Patient is within the FDA labeled age for the requested agent

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., dermatologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Zoryve Foam PA

## Drug Name(s)

Zoryve Foam

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

## **Exclusion Criteria:**

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of seborrheic dermatitis AND
- 2. ONE of the following:
  - A. Patient has tried and failed ONE topical antifungal (e.g., ketoconazole) OR ONE topical corticosteroid (e.g., betamethasone) OR
  - B. Patient has an intolerance or hypersensitivity to a topical antifungal OR a topical steroid OR
  - C. Patient has an FDA labeled contraindication to a topical antifungal OR a topical steroid

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of seborrheic dermatitis AND
- 3. Patient has had clinical benefit with the requested agent

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

## **Prescriber Restrictions:**

Prescriber is a specialist in the area of the patient's diagnosis (e.g., dermatologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## **Coverage Duration:**

Approval will be for 12 months

Ztalmy PA

## Drug Name(s)

Ztalmy

#### **Indications:**

All FDA-Approved Indications.

#### Off-Label Uses:

#### **Exclusion Criteria:**

## **Required Medical Information:**

Criteria for initial approval require BOTH of the following:

- 1. Patient has a diagnosis of seizures associated with cyclin-dependent kinase-like 5 (CDKL5) deficiency disorder (CDD) AND
- 2. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. BOTH of the following:
    - i. Patient's diagnosis has been confirmed with genetic testing indicating variant in CDKL5 gene AND
    - ii. Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

Criteria for renewal approval require ALL of the following:

- 1. Patient has been previously approved for the requested agent through the plan's Prior Authorization criteria AND
- 2. Patient has a diagnosis of seizures associated with cyclin-dependent kinase-like 5 (CDKL5) deficiency disorder (CDD) AND
- 3. ONE of the following:
  - A. There is evidence of a claim that the patient is currently being treated with the requested agent within the past 180 days OR
  - B. Prescriber states the patient is currently being treated with the requested agent OR
  - C. BOTH of the following:
    - i. Patient has had clinical benefit with the requested agent AND
    - ii. Prescriber is a specialist in the area of the patient's diagnosis (e.g., neurologist) or the prescriber has consulted with a specialist in the area of the patient's diagnosis

## Age Restriction:

Patient is within the FDA labeled age for the requested agent

**Prescriber Restrictions:** 

## **Coverage Duration:**

Approval will be for 12 months

Zyclara PA

## Drug Name(s)

Imiquimod (Zyclara)

**Imiquimod Pump** 

Zyclara

Zyclara Pump

## Indications:

All FDA-Approved Indications.

## Off-Label Uses:

**Exclusion Criteria:** 

## **Required Medical Information:**

Criteria for approval require ONE of the following:

- 1. Patient has a diagnosis of actinic keratosis OR
- 2. Patient has a diagnosis of external genital and/or perianal warts/condyloma acuminata AND the requested agent is Zyclara/imiquimod 3.75%

## Age Restriction:

#### **Prescriber Restrictions:**

## **Coverage Duration:**

External genital and/or perianal warts/condyloma acuminata: 8 weeks

Actinic keratosis: 6 weeks