South Dakota Department of Social Services

Medicaid P&T Committee Meeting December 2, 2022



Table of Contents

Agenda	2
Minutes	3
PA update	6
Top 15 Therapeutic Classes	9
Top 50 Drugs	10
Narrow Therapeutic Index drugs review	12
Opioid update	17
PA forms & criteria	22
Xelstyrm	155

DEPARTMENT OF SOCIAL SERVICES



DIVISION OF MEDICAL SERVICES 700 GOVERNORS DRIVE PIERRE, SD 57501-2291 PHONE: 605-773-3495 FAX: 605-773-5246

WEB: dss.sd.gov

SOUTH DAKOTA MEDICAID P&T COMMITTEE MEETING AGENDA

December 2, 2022 1:00 – 3:00 PM

Meeting Link:

https://teams.microsoft.com/l/meetupjoin/19%3ameeting OGU3NGFIZDEtZDhhMy00ZjYyLWI0ODgtYmRhMTMxODEwMTA0%40thread.v2/0?cont ext=%7b%22Tid%22%3a%22db05faca-c82a-4b9d-b9c5-0f64b6755421%22%2c%22Oid%22%3a%22b6efd724-b34e-4a86-b34c-e34f07dd4ceb%22%7d

Join with a video conferencing device

<u>425899727@t.plcm.vc</u> Video Conference ID: 111 219 836 59

Join by phone

+1 952-222-7450

Phone Conference ID: 113 036 048#

Call to order

Approval of previous meeting minutes

PA update

Review of top 15 therapeutic categories/top 50 drugs

Old business

Performance Measures Narrow Therapeutic Index (NTI) drugs Opioid update

New business

Biosimilar presentation Review PA forms & criteria Xelstyrm

Public input accepted after individual topic discussion Next meeting date March 24, 2023 & adjournment

South Dakota Department of Social Services, Division of Medicaid Services Pharmacy & Therapeutics (P&T) Committee Meeting Minutes

Friday, September 23, 2022 1:00 – 3:00 pm CT

Members and DSS Staff

Michelle Baack, MD	Х	Heather Preuss, MD	-
Dana Darger, RPh, Chair	Х	Matthew Stanley, DO	-
Mikel Holland, MD	Х	Deidre Van Gilder, PharmD	Х
Bill Ladwig, RPh	Х	Mike Jockheck, DSS Staff	Х
Kelley Oehlke, PharmD	Х	Matthew Ballard, DSS Staff	Х
Lenny Petrik, PharmD	-	Sarah Aker, DSS Staff	Х

Administrative Business

Darger called the meeting to order at 1:03 pm. The minutes of the June meeting were presented. Baack made a motion to approve. Van Gilder seconded the motion. The motion was unanimously approved.

Prior Authorization Update (PA) and Statistics

The committee reviewed the PA activity report from April 1, 2022, to June 30, 2022. A total of 1,791 PAs were reviewed of which 117 requests (6.5%) were received via telephone and 1,031 requests (57.6%) were received via fax, and 663 (35.9%) were reviewed via electronically. There was a 3.2% increase of PAs received compared to the previous quarter.

Analysis of the Top 15 Therapeutic Classes and Drug Spend

The committee reviewed the top 15 therapeutic classes by total cost of claims from April 1, 2022, to June 30, 2022. The top five therapeutic classes based on paid amount were atypical antipsychotics, disease-modifying anti-rheumatic agents, skin and mucous membrane agents, cystic fibrosis correctors, and hemostatics. These top 15 therapeutic classes make up 25.12 % of total claims. The committee also reviewed the top 50 drugs based on amount paid and number of claims. The top 50 drugs by amount paid make up 9.38 % of total claims. Of note, Opsumit made its debut on the top 50 drugs by paid amount. There was a comment regarding Eliquis starter kit. Darger requested to presentation on biosimilars and bioidenticals. Darger inquired if there was any public comment. There was none.

Old Business

Performance Measures

Samantha Moon from the Department of Medical Services provided follow up on two Performance Measures that the State is tracking: Care for Children Prescribed ADHD Medications: Ages 6-12 years old and Metabolic Monitoring for Children and Adolescents on Antipsychotics: Ages 1-17 years old. Committee discussed ways to ensure appropriate follow up care. Committee to discuss a possible PA renewal requirement at the next meeting. Darger inquired if there was any public comment. There was none.

Narrow Therapeutic Index Drugs

The committee reviewed the NTI utilization. Ladwig questioned the need for NTI list. Darger cited many states not having an NTI list anymore. After discussion, Ladwig made a motion to remove the NTI list.

Baack seconded the motion. Darger inquired if there was any public comment. There was none. The motion was approved unanimously. Van Gilder initiated discussion on adding PA to levothyroxine capsules. Ladwig made a motion to PA the capsule. Holland seconded the motion. Jockheck inquired about the criteria, for example, the trial and failure of a tablet before capsule is allowed over 180 days. Discussion ensued that most members would meet this requirement since most would have been on therapy for years. Committee will be provided more in-depth analysis especially how many claims are submitted with DAW 1 before removing the NTI drug list and PA on levothyroxine tablets.

Oseltamivir

The committee reviewed the NTI utilization. Jockheck reminded the committee oseltamivir's debut on the Top 50 drug list by paid amount last quarter. Committee commented utilization looked appropriate. Darger inquired if there was any public comment. There was none.

Xifaxan

Darger provided background information on the Xifaxan review. There is no utilization for Xifaxan 200mg due to a drug shortage, but three 200mg tablets are cheaper than one 550mg tablet. Since the diagnosis is coded directly with the drug strength, Van Gilder made a motion to remove specific strength to diagnosis. Baack seconded the motion. Darger inquired if there was any public comment. There was none. The motion was approved unanimously.

Sedative Hypnotics – doxepin

The committee reviewed doxepin utilization. No recommendation was given.

Vuity

The committee reviewed Vuity utilization. Baack recommended reviewing Vuity at the March 2023 meeting.

Opioid and muscle relaxant combination

The committee reviewed opioid utilization of members taking over 90 MME and members taking opioid in combination with muscle relaxants. No recommendation was given.

Opioid and stimulant

The committee reviewed opioid utilization of members taking over 90 MME and stimulant combination. No recommendation was given.

Opioid update

The committee reviewed 2Q2022 opioid outcomes compared to previous quarters from the opioid initiatives. The opioid figures for 2Q2022 excluded IHS utilization with the last similar comparison during 4Q2019. There was a decrease in opioid utilization and utilizers during 2Q2022 compared to 4Q2019 even with an increase in total eligible members.

Darger inquired if there was any public comment. There was none.

New Business

Flegsuvy

Fleqsuvy clinical information was presented for review. Baack recommended reviewing utilization at the March 2023 meeting. Darger inquired if there was any public comment. There were none.

Selgentis

Seglentis clinical information was presented for review. Committee recommended reviewing utilization at the March 2023 meeting. Darger inquired if there was any public comment. There were none.

Adjournment

The next meeting is scheduled on December 2, 2022. The March meeting is tentatively scheduled for March 24, 2022. The Committee made a motion to adjourn the meeting, and everyone seconded the motion. The motion passed unanimously, and the meeting adjourned at 2:52 pm.

PA Report 7/1/2022 – 9/30/2022

Compliance Summary

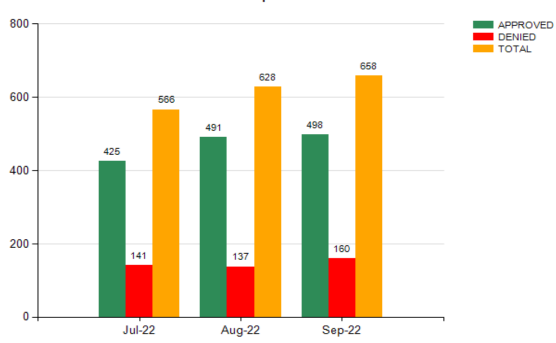
Priority	Total PAs	PAs Compliant	PAs Not Compliant	% PAs Compliant	% PAs Not Compliant
Standard	28	28	0	100.00%	0.00%
Urgent	1,824	1,824	0	100.00%	0.00%
Grand Total	1,852	1,852	0		

Drug Class	# of	Phone Requests		Fax Requests		Real-Time PA	
	Requests	#	%	#	%	#	%
Total	1,852	117	6.5%	1,031	57.6%	643	35.9%

PA Initial Requests Summary

Month	Approved	Denied	Total
Jul-22	425	141	566
Aug-22	491	137	628
Sep-22	498	160	658
3Q22	1,414	438	1,852
Percent of Total	76.35%	23.65%	

PA Requests Details



Top Therapeutic Classes for PA

Drug Class	Approved	Denied	Total	Approval Rate	% of Total Requests	Most Requested Products
ANTIPSYCHOTIC/ANTIMANIC	305	16	321	95.02%	17.33%	, INVEGA SUSTENNA
ANTIDIABETICS*	246	48	294	83.67%	15.87%	, OZEMPIC
DERMATOLOGICALS*	98	99	197	49.75%	10.64%	DUPIXENT, SPINOSAD
ANALGESICS - OPIOID*	101	83	184	54.89%	9.94%	TRAMADOL, HYDROCODONE
ANTIDEPRESSANTS*	153	12	165	92.73%	8.91%	, SERTRALINE
OTHERS -	511	180	691	73.95%	37.31%	
3Q22	1,414	438	1,852	76.35%		

PA Drug Class Summary

Drug Class	Approved	Denied	Total	Approval Rate
59 - ANTIPSYCHOTICS/ANTIMANIC AGENTS*	305	16	321	95.02%
27 - ANTIDIABETICS*	246	48	294	83.67%
90 - DERMATOLOGICALS*	98	99	197	49.75%
65 - ANALGESICS - OPIOID*	101	83	184	54.89%
58 - ANTIDEPRESSANTS*	153	12	165	92.73%
67 - MIGRAINE PRODUCTS*	59	42	101	58.42%
49 - ULCER DRUGS/ANTISPASMODICS/ANTICHOLINERG	74	12	86	86.05%
52 - GASTROINTESTINAL AGENTS - MISC.*	73	5	78	93.59%
61 - ADHD/ANTI-NARCOLEPSY/ANTI-OBESITY/ANOREX	42	19	61	68.85%
66 - ANALGESICS - ANTI-INFLAMMATORY*	45	12	57	78.95%
41 - ANTIHISTAMINES*	30	10	40	75.00%
16 - ANTI-INFECTIVE AGENTS - MISC.*	31	3	34	91.18%
72 - ANTICONVULSANTS*	29	4	33	87.88%
12 - ANTIVIRALS*	4	21	25	16.00%
54 - URINARY ANTISPASMODICS*	21	4	25	84.00%
30 - ENDOCRINE AND METABOLIC AGENTS - MISC.*	10	10	20	50.00%
39 - ANTIHYPERLIPIDEMICS*	6	6	12	50.00%
62 - PSYCHOTHERAPEUTIC AND NEUROLOGICAL AGENT	12	0	12	100.00%
75 - MUSCULOSKELETAL THERAPY AGENTS*	7	5	12	58.33%
21 - ANTINEOPLASTICS AND ADJUNCTIVE THERAPIES	11	0	11	100.00%
83 - ANTICOAGULANTS*	7	4	11	63.64%
33 - BETA BLOCKERS*	7	1	8	87.50%
02 - CEPHALOSPORINS*	3	4	7	42.86%
50 - ANTIEMETICS*	4	3	7	57.14%
03 - MACROLIDES*	4	2	6	66.67%
44 - ANTIASTHMATIC AND BRONCHODILATOR AGENTS*	5	1	6	83.33%
40 - CARDIOVASCULAR AGENTS - MISC.*	3	2	5	60.00%
34 - CALCIUM CHANNEL BLOCKERS*	3	1	4	75.00%
42 - NASAL AGENTS - SYSTEMIC AND TOPICAL*	2	2	4	50.00%
19 - PASSIVE IMMUNIZING AND TREATMENT AGENTS*	3	0	3	100.00%
36 - ANTIHYPERTENSIVES*	3	0	3	100.00%
45 - RESPIRATORY AGENTS - MISC.*	1	2	3	33.33%
60 - HYPNOTICS/SEDATIVES/SLEEP DISORDER AGENT	2	1	3	66.67%
57 - ANTIANXIETY AGENTS*	2	0	2	100.00%
82 - HEMATOPOIETIC AGENTS*	2	0	2	100.00%
86 - OPHTHALMIC AGENTS*	0	2	2	0.00%
99 - MISCELLANEOUS THERAPEUTIC CLASSES*	2	0	2	100.00%
01 - PENICILLINS*	0	1	1	0.00%
32 - ANTIANGINAL AGENTS*	1	0	<u>+</u> 1	100.00%
56 - GENITOURINARY AGENTS - MISCELLANEOUS*	0	1	1	0.00%
79 - MINERALS & ELECTROLYTES*	1	0	1	100.00%
80 - NUTRIENTS*	1	0	1	100.00%
93 - ANTIDOTES AND SPECIFIC ANTAGONISTS*	1	0	1	100.00%
3Q22	1,414	438	1,852	100.0070
		730	1.032	

PA Appeals Summary

Month	Approved	Approved %	Denied	Denied %	Total
Jul-22	11	61.11%	7	38.89%	18
Aug-22	16	69.57%	7	30.43%	23
Sep-22	15	68.18%	7	31.82%	22
3Q22	42	66.67%	21	33.33%	63

Appeals Detail

Drug Class	Approved	Denied	Total	Approval Rate
DUPIXENT	5	1	6	83.33%
AIMOVIG	4	0	4	100.00%
MAVYRET	2	8	10	20.00%
LUBIPROSTONE	2	1	3	66.67%
NURTEC	2	1	3	66.67%
SPINOSAD	2	1	3	66.67%
STELARA	2	1	3	66.67%
EMGALITY	2	0	2	100.00%
IVERMECTIN	2	0	2	100.00%
TRAMADOL HCL	2	0	2	100.00%
OZEMPIC	1	3	4	25.00%
AJOVY	1	0	1	100.00%
AMPHETAMINE/DEXTROAMPHETAMINE	1	0	1	100.00%
CEPHALEXIN	1	0	1	100.00%
EPIDIOLEX	1	0	1	100.00%
ESZOPICLONE	1	0	1	100.00%
HYDROCODONE BITARTRATE/APAP	1	0	1	100.00%
INGREZZA	1	0	1	100.00%
JYNARQUE	1	0	1	100.00%
MALATHION	1	0	1	100.00%
METRONIDAZOLE	1	0	1	100.00%
NORDITROPIN FLEXPRO	1	0	1	100.00%
OPZELURA	1	0	1	100.00%
OTEZLA	1	0	1	100.00%
PALYNZIQ	1	0	1	100.00%
SOFOSBUVIR/VELPATASVIR	1	0	1	100.00%
UBRELVY	1	0	1	100.00%
CABERGOLINE	0	1	1	0.00%
EPCLUSA	0	1	1	0.00%
ESOMEPRAZOLE MAGNESIUM	0	1	1	0.00%
HUMIRA PEN-PS/UV STARTER	0	1	1	0.00%
LINZESS	0	1	1	0.00%
3Q22	42	21	63	

Top 15 Therapeutic Classes & Top 50 Drugs

T	TOP 15 THERAPEUTIC CLASSES BASED ON NUMBER OF CLAIMS FROM 7/1/2022 – 9/30/2022							
	AHFS Description	Total Rxs	Plan Paid Amount	Paid/Rx	% Total Claims			
1	SELECTIVE-SEROTONIN REUPTAKE INHIBITORS	15,214	\$195,845.45	\$12.87	6.76%			
2	ANTICONVULSANTS, MISCELLANEOUS	12,154	\$954,170.14	\$78.51	5.40%			
3	ATYPICAL ANTIPSYCHOTICS	9,569	\$3,088,251.45	\$322.74	4.25%			
4	SELECTIVE BETA-2-ADRENERGIC AGONISTS	8,147	\$493,705.47	\$60.60	3.62%			
5	SECOND GENERATION ANTIHISTAMINES	8,048	\$91,911.48	\$11.42	3.58%			
6	RESPIRATORY AND CNS STIMULANTS	7,393	\$517,725.26	\$70.03	3.28%			
7	AMPHETAMINES	7,298	\$1,259,543.69	\$172.59	3.24%			
8	PROTON-PUMP INHIBITORS	6,637	\$187,143.51	\$28.20	2.95%			
9	ADRENALS	6,406	\$687,083.15	\$107.26	2.85%			
10	AMINOPENICILLIN ANTIBIOTICS	6,367	\$93,022.19	\$14.61	2.83%			
11	OPIATE AGONISTS	5,739	\$178,532.18	\$31.11	2.55%			
12	ANXIOLYTICS, SEDATIVES, & HYPNOTICS, MISC	5,171	\$69,370.32	\$13.42	2.30%			
13	CONTRACEPTIVES	4,168	\$131,564.27	\$31.57	1.85%			
14	CENTRAL NERVOUS SYSTEM AGENTS, MISC.	4,135	\$226,136.79	\$54.69	1.84%			
15	THYROID AGENTS	3,813	\$70,063.95	\$18.38	1.69%			
Tot	al	110,259	\$8,244,069.30	\$74.77	48.98%			

	TOP 15 THERAPEUTIC CLASSES BASED ON AMOUNT PAID FROM 7/1/2022 – 9/30/2022								
	AHFS Description	Total Rxs	Plan Paid Amount	Paid/Rx	% Total Claims				
1	ATYPICAL ANTIPSYCHOTICS	9,569	\$3,088,251.45	\$322.74	4.25%				
2	DISEASE-MODIFYING ANTIRHEUMATIC AGENTS	397	\$2,636,653.65	\$6,641.44	0.18%				
3	SKIN AND MUCOUS MEMBRANE AGENTS, MISC.	745	\$2,170,886.51	\$2,913.94	0.33%				
4	CYSTIC FIBROSIS (CFTR) CORRECTORS	69	\$1,561,239.48	\$22,626.66	0.03%				
5	AMPHETAMINES	7,298	\$1,259,543.69	\$172.59	3.24%				
6	HEMOSTATICS	64	\$1,097,475.55	\$17,148.06	0.03%				
7	ANTICONVULSANTS, MISCELLANEOUS	12,154	\$954,170.14	\$78.51	5.40%				
8	INCRETIN MIMETICS	1,083	\$922,322.09	\$851.64	0.48%				
9	ANTINEOPLASTIC AGENTS	296	\$842,309.68	\$2,845.64	0.13%				
10	ADRENALS	6,406	\$687,083.15	\$107.26	2.85%				
11	LONG-ACTING INSULINS	1,439	\$636,346.45	\$442.21	0.64%				
12	RAPID-ACTING INSULINS	1,391	\$563,131.83	\$404.84	0.62%				
13	GI DRUGS, MISCELLANEOUS	435	\$547,557.45	\$1,258.75	0.19%				
14	RESPIRATORY AND CNS STIMULANTS	7,393	\$517,725.26	\$70.03	3.28%				
15	SELECTIVE BETA-2-ADRENERGIC AGONISTS	8,147	\$493,705.47	\$60.60	3.62%				
Tot	al	56,886	\$17,978,401.85	\$316.04	25.27%				

Total Rx Claims from 7/1/2022 – 9/30/2022	225,090
---	---------

	TOP 50 DRUGS BASED ON NUMBER OF CLAIMS FROM 7/1/2022 – 9/30/2022							
	AHFS Description	Drug Label Name	Total Rxs	Plan Paid Amount	Paid/Rx	% Total Claims		
1	SELECTIVE-SEROTONIN REUPTAKE INHIBITORS	FLUOXETINE HCL	5,387	\$67,869.77	\$12.60	2.39%		
2	RESPIRATORY AND CNS STIMULANTS	METHYLPHENIDATE HCL	5,151	\$228,975.97	\$44.45	2.29%		
3	SELECTIVE-SEROTONIN REUPTAKE INHIBITORS	SERTRALINE HCL	4,819	\$57,394.85	\$11.91	2.14%		
4	AMINOPENICILLIN ANTIBIOTICS	AMOXICILLIN	4,697	\$60,557.50	\$12.89	2.09%		
5	SELECTIVE BETA-2-ADRENERGIC AGONISTS	ALBUTEROL SULFATE HFA	4,653	\$179,702.43	\$38.62	2.07%		
6	SECOND GENERATION ANTIHISTAMINES	CETIRIZINE HCL	4,615	\$49,423.02	\$10.71	2.05%		
7	PROTON-PUMP INHIBITORS	OMEPRAZOLE	3,936	\$45,208.69	\$11.49	1.75%		
8	AMPHETAMINES	VYVANSE	3,615	\$1,140,989.36	\$315.63	1.61%		
9	SELECTIVE-SEROTONIN REUPTAKE INHIBITORS	ESCITALOPRAM OXALATE	3,527	\$44,724.97	\$12.68	1.57%		
10	ANTICONVULSANTS, MISCELLANEOUS	GABAPENTIN	3,509	\$57,293.43	\$16.33	1.56%		
11	LEUKOTRIENE MODIFIERS	MONTELUKAST SODIUM	3,505	\$45,567.05	\$13.00	1.56%		
12	AMPHETAMINES	AMPHETAMINE/DEXTROAM	3,494	\$90,463.86	\$25.89	1.55%		
13	SEROTONIN MODULATORS	TRAZODONE HCL	3,474	\$35,047.41	\$10.09	1.54%		
14	THYROID AGENTS	LEVOTHYROXINE SODIUM	3,071	\$43,958.18	\$14.31	1.36%		
15	CENTRAL ALPHA-AGONISTS	CLONIDINE HCL	2,841	\$35,549.78	\$12.51	1.26%		
16	ANTIDEPRESSANTS, MISCELLANEOUS	BUPROPION HCL	2,639	\$50,545.16	\$19.15	1.17%		
17↑	BIGUANIDES	METFORMIN HCL	2,515	\$32,484.28	\$12.92	1.12%		
18	OPIATE AGONISTS	HYDROCODONE BITARTR/AC	2,322	\$33,878.71	\$14.59	1.03%		
19	ATYPICAL ANTIPSYCHOTICS	ARIPIPRAZOLE	2,235	\$31,648.39	\$14.16	0.99%		
20	HMG-COA REDUCTASE INHIBITORS	ATORVASTATIN CALCIUM	2,229	\$26,032.62	\$11.68	0.99%		
21	ANGIOTENSIN-CONVERTING ENZYME INHIBIT	LISINOPRIL	2,193	\$21,517.54	\$9.81	0.97%		
22	CENTRAL NERVOUS SYSTEM AGENTS, MISC	GUANFACINE ER	2,068	\$35,921.71	\$17.37	0.92%		
23	SEL.SEROTONIN, NOREPI REUPTAKE INHIBIT	DULOXETINE HCL	2,007	\$30,958.39	\$15.43	0.89%		
24	1ST GENERATION CEPHALOSPORIN ANTIBIOT	CEPHALEXIN	1,998	\$31,894.17	\$15.96	0.89%		
25	ANXIOLYTICS, SEDATIVES, & HYPNOTICS, MISC	HYDROXYZINE HCL	1,941	\$24,437.97	\$12.59	0.86%		
26	ANTICONVULSANTS, MISCELLANEOUS	LAMOTRIGINE	1,847	\$25,393.24	\$13.75	0.82%		
27	ATYPICAL ANTIPSYCHOTICS	RISPERIDONE	1,844	\$22,033.63	\$11.95	0.82%		
28	SECOND GENERATION ANTIHISTAMINES	LORATADINE	1,805	\$19,811.36	\$10.98	0.80%		
29	ADRENALS	PREDNISONE	1,746	\$17,231.13	\$9.87	0.78%		
30↑	SELECTIVE BETA-2-ADRENERGIC AGONISTS	ALBUTEROL SULFATE	1,723	\$31,249.39	\$18.14	0.77%		
31	AMINOPENICILLIN ANTIBIOTICS	AMOXICILLIN/CLAVULANATE	1,665	\$31,889.23	\$19.15	0.74%		
32	ATYPICAL ANTIPSYCHOTICS	QUETIAPINE FUMARATE	1,626	\$20,047.57	\$12.33	0.72%		
33	5-HT3 RECEPTOR ANTAGONISTS	ONDANSETRON ODT	1,587	\$23,349.38	\$14.71	0.71%		
34	ANXIOLYTICS, SEDATIVES, & HYPNOTICS, MISC	BUSPIRONE HCL	1,585	\$20,318.06	\$12.82	0.70%		
35	CORTICOSTEROIDS (EENT)	FLUTICASONE PROPIONATE	1,568	\$22,833.56	\$14.56	0.70%		
36	BENZODIAZEPINES (ANTICONVULSANTS)	CLONAZEPAM	1,533	\$17,068.72	\$11.13	0.68%		
37	CORTICOSTEROID-SKIN, MUCOUS MEMBRAN	TRIAMCINOLONE ACETONID	1,531	\$23,370.54	\$15.26	0.68%		
38	OTHER MACROLIDE ANTIBIOTICS	AZITHROMYCIN	1,512	\$23,740.14	\$15.70	0.67%		
39	COMPOUNDS	-	1,473	\$38,052.67	\$25.83	0.65%		
40	CENTRALLY ACTING SKELETAL MUSCLE RELAX	CYCLOBENZAPRINE HCL	1,344	\$13,563.11	\$10.09	0.60%		
41	ANTICONVULSANTS, MISCELLANEOUS	LEVETIRACETAM	1,330	\$27,648.96	\$20.79	0.59%		
42	3RD GENERATION CEPHALOSPORIN ANTIBIO	CEFDINIR	1,320	\$25,543.80	\$19.35	0.59%		
43	PROTON-PUMP INHIBITORS	PANTOPRAZOLE SODIUM	1,311	\$18,249.72	\$13.92	0.58%		
44	ANTICONVULSANTS, MISCELLANEOUS	TOPIRAMATE	1,305	\$16,755.35	\$12.84	0.58%		
45	DIHYDROPYRIDINES	AMLODIPINE BESYLATE	1,272	\$12,506.70	\$9.83	0.57%		
46↑	ANTIBACTERIALS (SKIN, MUCOUS MEMBRAN)	MUPIROCIN	1,261	\$20,931.20	\$16.60	0.56%		
47	ANGIOTENSIN II RECEPTOR ANTAGONISTS	LOSARTAN POTASSIUM	1,149	\$13,726.03	\$11.95	0.51%		
48	VITAMIN B COMPLEX	FOLIC ACID	1,134	\$10,040.96	\$8.85	0.50%		
49	ANTIDEPRESSANTS, MISCELLANEOUS	MIRTAZAPINE	1,127	\$15,883.20	\$14.09	0.50%		
50	VITAMIN D	VITAMIN D	1,125	\$11,278.18	\$10.03	0.50%		
	Total Top 50 Drugs		119,164	\$3,024,561.04	\$25.38	52.94%		
			- , -		*			

	TOP 50 DRUGS BASED ON AMOUNT PAID FROM 7/1/2022 – 9/30/2022								
	AHFS Description	Drug Label Name	Total Rxs	Plan Paid Amount	Paid/Rx	% Total Claims			
1	DISEASE-MODIFYING ANTIRHEUMATIC AGT	HUMIRA & PEN	164	\$1,429,002.36	\$8,713.43	0.07%			
2	CYSTIC FIBROSIS (CFTR) CORRECTORS	TRIKAFTA	51	\$1,230,589.71	\$24,129.21	0.02%			
3	AMPHETAMINES	VYVANSE	3,615	\$1,140,989.36	\$315.63	1.61%			
4	ATYPICAL ANTIPSYCHOTICS	INVEGA TRNZA/SUSTNA/HAFYRA	352	\$973,889.67	\$2,766.73	0.16%			
5	SKIN & MUCOUS MEMBRANE AGENTS	STELARA	41	\$925,977.81	\$22,584.82	0.02%			
6	SKIN & MUCOUS MEMBRANE AGENTS	DUPIXENT	224	\$752,467.11	\$3,359.23	0.10%			
7	ATYPICAL ANTIPSYCHOTICS	LATUDA	450	\$576,653.61	\$1,281.45	0.20%			
8	INCRETIN MIMETICS	OZEMPIC	614	\$520,774.40	\$848.17	0.27%			
9	ATYPICAL ANTIPSYCHOTICS	ARISTADA & INTIO	158	\$428,414.56	\$2,711.48	0.07%			
10	ATYPICAL ANTIPSYCHOTICS	VRAYLAR	327	\$380,709.78	\$1,164.25	0.15%			
11	DISEASE-MODIFYING ANTIRHEUMATIC AGT	COSENTYX & SENSOREADY	49	\$333,382.27	\$6,803.72	0.02%			
12	CYSTIC FIBROSIS (CFTR) CORRECTORS	ORKAMBI	18	\$330,649.77	\$18,369.43	0.01%			
13	DISEASE-MODIFYING ANTIRHEUMATIC AGT	ENBREL, MINI, SURECLICK	53	\$328,959.28	\$6,206.78	0.02%			
14	SOMATOTROPIN AGONISTS	NORDITROPIN FLEXPRO	81	\$311,715.05	\$3,848.33	0.04%			
15	HEMOSTATICS	HEMLIBRA	6	\$303,221.52	\$50,536.92	0.00%			
16	ANTICONVULSANTS, MISCELLANEOUS	EPIDIOLEX	117	\$300,075.87	\$2,564.75	0.05%			
17	SODIUM-GLUC COTRANSPORT 2 INHIBITOR	JARDIANCE	527	\$273,753.54	\$519.46	0.23%			
18	ATYPICAL ANTIPSYCHOTICS	REXULTI	225	\$272,791.47	\$1,212.41	0.10%			
19	MUCOLYTIC AGENTS	PULMOZYME	60	\$249,672.08	\$4,161.20	0.03%			
20↑	SKIN & MUCOUS MEMBRANE AGENTS	TALTZ	33	\$246,123.33	\$7,458.28	0.01%			
21	RESPIRATORY AND CNS STIMULANTS	METHYLPHENIDATE HCL	5,151	\$228,975.97	\$44.45	2.29%			
22	INCRETIN MIMETICS	TRULICITY	264	\$228,876.84	\$866.96	0.12%			
23	GI DRUGS, MISCELLANEOUS	GATTEX	5	\$214,620.00	\$42,924.00	0.00%			
24	HEMOSTATICS	ADVATE	12	\$211,890.76	\$17,657.56	0.01%			
25	LONG-ACTING INSULINS	LANTUS & SOLOSTAR	535	\$208,725.86	\$390.14	0.24%			
26	HIV INTEGRASE INHIBITOR ANTIRETROVIRA	BIKTARVY	59	\$202,378.73	\$3,430.15	0.03%			
27	RIFAMYCIN ANTIBIOTICS	XIFAXAN	79	\$193,996.17	\$2,455.65	0.04%			
28	OTHER MISCELLANEOUS THERAPEUTIC AGT	EVRYSDI	8	\$187,854.80	\$23,481.85	0.00%			
29↑	SELECTIVE BETA-2-ADRENERGIC AGONISTS	ALBUTEROL SULFATE HFA	4,653	\$179,702.43	\$38.62	2.07%			
30	VESICULAR MONOAMINE TRANSPORT2 INH	INGREZZA	25	\$175,608.05	\$7,024.32	0.01%			
31	LONG-ACTING INSULINS	TRESIBA FLEXTOUCH	310	\$167,185.68	\$539.31	0.14%			
32↑	SKIN & MUCOUS MEMBRANE AGENTS,	TREMFYA	13	\$163,680.37	\$12,590.80	0.01%			
33	HEMOSTATICS	RECOMBINATE	3	\$150,811.35	\$50,270.45	0.00%			
34↑	DIRECT FACTOR XA INHIBITORS	ELIQUIS & STARTER	317	\$149,629.18	\$472.02	0.14%			
35	HEMOSTATICS	NOVOSEVEN RT	2	\$148,821.10	\$74,410.55	0.00%			
36	ATYPICAL ANTIPSYCHOTICS	ABILIFY MAINTENA	60	\$148,383.06	\$2,473.05	0.03%			
37	SKIN & MUCOUS MEMBRANE AGENTS	SKYRIZI & PEN	8	\$145,372.44	\$18,171.56	0.00%			
38	ADRENALS	FLOVENT HFA	614	\$144,669.50	\$235.62	0.27%			
39	SELECTIVE BETA-2-ADRENERGIC AGONISTS	ADVAIR HFA	401	\$144,487.40	\$360.32	0.18%			
40	LONG-ACTING INSULINS	LEVEMIR & FLEXTOUCH	299	\$141,731.05	\$474.02	0.13%			
41	RAPID-ACTING INSULINS	INSULIN ASPART FLEXPEN	384	\$132,582.71	\$345.27	0.17%			
42	HIV INTEGRASE INHIBITOR ANTIRETROVIRA	GENVOYA	37	\$129,334.04	\$3,495.51	0.02%			
43	HEMOSTATICS	XYNTHA SOLOFUSE	3	\$127,883.55	\$42,627.85	0.00%			
44	GI DRUGS, MISCELLANEOUS	CHOLBAM	6	\$124,413.30	\$20,735.55	0.00%			
45↓	ENZYMES	PALYNZIQ	3	\$117,841.65	\$39,280.55	0.00%			
46↑	GI DRUGS, MISCELLANEOUS	LINZESS	252	\$115,447.61	\$458.13	0.11%			
47↑	ALPHA- AND BETA-ADRENERGIC AGONISTS	EPINEPHRINE	397	\$114,384.01	\$288.12	0.18%			
48	DIPEPTIDYL PEPTIDASE-4(DPP-4) INHIBITOR	JANUVIA	232	\$111,219.03	\$479.39	0.10%			
49	RAPID-ACTING INSULINS	NOVOLOG FLEXPEN	184	\$107,141.13	\$582.29	0.08%			
50	ANTIMUSCARINICS/ANTISPASMODICS	SPIRIVA RESPIMAT	223	\$103,317.28	\$463.31	0.10%			

Old Business

Performance Measures

Narrow Therapeutic Index (NTI) Drugs

FDA US Food & Drug Administration FY2015 Regulatory Science Research Report: Narrow Therapeutic Index Drugs: Narrow therapeutic index drugs are drugs where small differences in dose or blood concentration may lead to serious therapeutic failures and/or adverse drug reactions that are lifethreatening or result in persistent or significant disability or incapacity.

South Dakota NTI drug list

Therapeutic Class

carbamazepine

cyclosporine

digoxin

lamotrigine

levetiracetam

lithium

• Pancreatic Drug Products

phenytoin

• procainamide

• quinidine

thyroid preparations

theophylline

topiramatevalproic Acid

• warfarin

Example Brand Names:

Tegretol

Neoral, Sandimmune

Lanoxin, Digitek

Lamictal/XR

Keppra

Lithobid, Eskalith

Creon, Pancreaze

Dilantin, Phenytek

Pronestyl, Procanbid

Quinidex, Quinaglute, Quinamm

Synthroid, Levothroid, Armour Thyroid

Aminophylline, Elixophyllin, Theo-24, Theo-Dur,

Theo-chron, Uniphyl

Topamax

Depakene

Coumadin, Jantoven

Other States' NTI drug list:

State A

- Coumadin
- Dilantin
- Lanoxin
- Premarin
- Provera
- Synthroid
- Tegretol

State B

- Dilantin
- Tegretol

Dispense As Written Definitions

CODE	DESCRIPTION
	No Product Selection Indicated - This is the field default value that is appropriately used for prescriptions for
0	single source brand, co-branded/co-licensed, or generic products. For a multi-source branded product with
	available generic(s), DAW 0 is not appropriate, and may result in a reject.
	Substitution Not Allowed by Prescriber – This value is used when the prescriber indicates, in a manner
1	specified by prevailing law, that the product is Medically Necessary to be Dispensed As Written. DAW 1 is
	based on prescriber instruction and not product classification.
	<u>Substitution Allowed-Patient Requested Product Dispensed</u> -This value is used when the prescriber has
2	indicated, in a manner specified by prevailing law, that generic substitution is permitted and the patient
-	requests the brand product. This situation can occur when the prescriber writes the prescription using either
	the brand or generic name and the product is available from multiple sources.
	Substitution Allowed-Pharmacist Selected Product Dispensed-This value is used when the prescriber has
3	indicated, in a manner specified by prevailing law, that generic substitution is permitted and the pharmacist
	determines that the brand product should be dispensed. This can occur when the prescriber writes the
	prescription using either the brand or generic name and the product is available from multiple sources.
	Substitution Allowed-Generic Drug Not in Stock-This value is used when the prescriber has indicated, in a
4	manner specified by prevailing law, that generic substitution is permitted and the brand product is dispensed
'	since a currently marketed generic is not stocked in the pharmacy. This situation exists due to the buying
	habits of the pharmacist, not because of the unavailability of the generic product in the marketplace.
	<u>Substitution Allowed-Brand Drug Dispensed as a Generic</u> -This value is used when the prescriber has indicated,
5	in a manner specified by prevailing law, that generic substitution is permitted and the pharmacist is utilizing
	the brand product as the generic entity.
6	Override-This value is used by various claims processors in very specific instances as defined by that claims'
	processor and/or its client(s).
	<u>Substitution Not Allowed-Brand Drug Mandated by Law</u> -This value is used when the prescriber has indicated,
7	in a manner specified by prevailing law, that generic substitution is permitted but prevailing law or regulation
-	prohibits the substitution of a brand product even though generic versions of the product may be available in
	the marketplace.
_	Substitution Allowed-Generic Drug Not Available in Marketplace-This value is used when the prescriber has
8	indicated, in a manner specified by prevailing law, that generic substitution is permitted and the brand product
	is dispensed since the generic is not currently manufactured, distributed, or is temporarily unavailable.
	Substitution Allowed By Prescriber but Plan Requests Brand - Patient's Plan Requested Brand Product To Be
_	<u>Dispensed</u> - This value is used when the prescriber has indicated, in a manner specified by prevailing law, that
9	generic substitution is permitted, but the plan's formulary requests the brand product. This situation can occur
	when the prescriber writes the prescription using either the brand or generic name and the product is available
	from multiple sources.

NTI Utilization

Time frame: 7/1/2022 to 10/31/2022

Carbamazepine	Total Rx	Paid Amount	Paid/Rx	Utilizers	Age Range
carbamazepine SUS • DAW 0 = 8	38	\$2,983.17	\$78.50	8	14 – 44
carbamazepine CHW • DAW 0 = 16	51	\$2,579.09	\$50.57	16	9 – 59
carbamazepine CAP ER • DAW 0 = 26	110	\$10,815.60	\$98.32	26	14 – 65
carbamazepine TAB ER • DAW 0 = 22	76	\$5,552.44	\$73.06	22	24 – 67
carbamazepine TAB • DAW 0 = 42	138	\$3,809.33	\$27.60	42	11 – 64
Epitol TAB • DAW 0 = 2	5	\$111.71	\$22.34	2	43, 49
TEGRETOL-XR TAB (MSB) • DAW 1 = 1	4	\$281.81	\$70.45	1	26

^{*}Red font denotes brand utilization

Lamotrigine	Total Rx	Paid Amount	Paid/Rx	Utilizers	Age Range
lamotrigine TAB • DAW 0 = 704	2,433	\$33,056.32	\$13.59	704	6 – 68
LAMICTAL TAB (MSB) ■ DAW 1 = 5	29	\$42,262.85	\$1,457.34	5	27 – 54
lamotrigine TAB ER • DAW 0 = 46	144	\$12,401.70	\$86.12	46	13 – 60
LAMICTAL TAB XR (MSB) • DAW 1 = 4	11	\$27,808.22	\$2,528.02	4	26 – 36
lamotrigine CHW • DAW 0 = 16	63	\$3,572.99	\$56.72	16	6 – 48
lamotrigine TAB ODT ■ DAW 0 = 6	13	\$5,112.92	\$393.30	6	6 – 23

^{*}Red font denotes brand utilization

Phenytoin	Total Rx	Paid Amount	Paid/Rx	Utilizers	Age Range
phenytoin CHW • DAW 0 = 6	26	\$830.12	\$31.93	6	9 – 63
DILANTIN CHW (MSB) • DAW 1 = 2	7	\$1,282.78	\$183.25	2	9, 32
DILANTIN 30 MG CAP (SSB) • DAW 0 = 3	7	\$925.67	\$132.24	3	33, 36
 phenytoin 100mg or 300mg CAP DAW 0 = 37 DAW 1 = 1 	127	\$3,596.39	\$28.32	38	31 – 64
phenytoin SUS • DAW 0 = 3	13	\$427.33	\$32.87	3	34 – 64

^{*}Red font denotes brand utilization

Levetiracetam	Total Rx	Paid Amount	Paid/Rx	Utilizers	Age Range
levetiracetam SOL DAW 0 = 170 DAW 1 = 1	638	\$13,685.97	\$21.45	171	0 – 62
KEPPRA SOL (MSB) • DAW 1 = 2	9	\$16,488.84	\$1,832.09	2	18, 21
levetiracetam TAB • DAW 0 = 368 • DAW 2 = 1	1,150	\$23,637.79	\$20.55	369	4 – 88
KEPPRA TAB (MSB) ■ DAW 1 = 2	11	\$5,458.41	\$496.22	2	34, 53
SPRITAM TAB (SSB) • DAW 0 = 1	4	\$2,446.12	\$611.53	1	12
levetiracetam TAB ER • DAW 0 = 21	65	\$2,243.60	\$34.52	21	13 – 61
KEPPRA TAB XR (MSB) • DAW = 1	4	\$5,708.08	\$1,427.02	1	23

Topiramate	Total Rx	Paid Amount	Paid/Rx	Utilizers	Age Range
topiramate TAB • DAW 0 = 573	1,721	\$20,957.59	\$12.18	573	22 – 66
TOPAMAX TAB (MSB) • DAW 1 = 1	8	\$6,266.92	\$783.37	1	26
topiramate CAP • DAW 0 = 10	24	\$1,532.88	\$63.87	10	5 – 43
topiramate CAP ER • DAW 0 = 10	20	\$5,893.43	\$294.67	10	12 – 56
TROKENDI XR CAP (SSB) • DAW 0 = 8	25	\$24,179.54	\$967.18	8	15 – 50
EPRONTIA SOL (SSB) • DAW 0 = 1	3	\$2,026.65	\$675.55	1	8

Valproic Acid	Total Rx	Paid Amount	Paid/Rx	Utilizers	Age Range
valproic acid CAP and SOL • DAW 0 = 62	363	\$8,521.74	\$23.48	83	0 – 63
divalproex CAP • DAW 0 = 43	155	\$8,452.72	\$54.54	43	3 – 63
divalproex TAB DR • DAW 0 = 159	651	\$11,371.80	\$17.47	159	4 – 75
DEPAKOTE TAB DR (MSB) • DAW 0 = 1	4	\$1,644.60	\$411.15	1	54
divalproex TAB ER • DAW 0 = 160	588	\$13,200.21	\$22.45	160	7 – 78
 DEPAKOTE ER TAB (MSB) DAW 1 = 3 DAW 2 = 1 	12	\$8,290.51	\$690.88	4	15 – 59
DEPAKOTE SPR CAP (MSB) • DAW 0 = 1 • DAW 1 = 3	15	\$5,007.94	\$690.88	4	18 – 37

^{*}Red font denotes brand utilization

Cyclosporine	Total Rx	Paid Amount	Paid/Rx	Utilizers	Age Range
cyclosporine CAP					
• DAW 0 = 4	17	\$1,979.88	\$116.46	6	16 – 43
• DAW 1 = 2					
NEORAL SOL 100MG/ML (MSB)	2	\$854.12	\$427.06	1	12
• DAW 0 = 1	2	\$654.12	\$427.00	1	12

^{*}Red font denotes brand utilization

Theophylline	Total Rx	Paid Amount	Paid/Rx	Utilizers	Age Range
theophylline TAB ER • DAW 0 = 3	8	\$128.27	\$16.03	3	40 – 53
THEO-24 CAP CR (SSB) • DAW 0 = 2	5	\$540.34	\$108.07	2	21 – 61

^{*}Red font denotes brand utilization

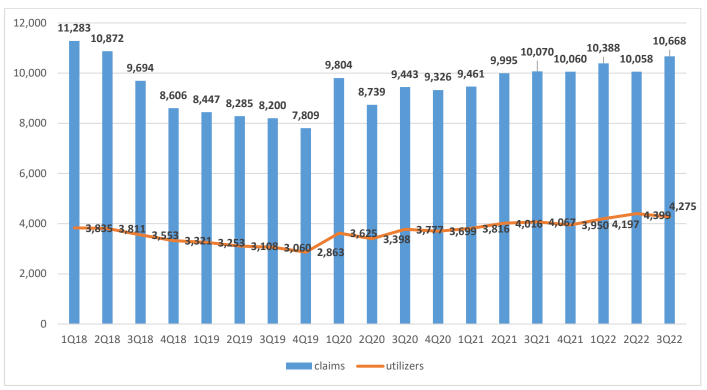
Thyroid Preparations	Total Rx	Paid Amount	Paid/Rx	Utilizers	Age Range
NP Thyroid TAB • DAW 0 = 29	82	\$4,042.81	\$49.30	29	10 – 63
 ARMOUR THYROID TAB (SSB) DAW 0 = 19 DAW 1 = 2 DAW 2 = 6 DAW 8 = 1 	109	\$4,832.30	\$44.33	28	9 – 63

^{*}Red font denotes brand utilization

Levothyroxine	Total Rx	Paid Amount	Paid/Rx	Utilizers	Age Range
levothyroxine CAP • DAW 0 = 12	28	\$3,110.45	\$111.09	12	15 – 49
levothyroxine TAB	4,147	\$55,613.03	\$13.41	1,361	0 – 88
Levoxyl TAB • DAW 1 = 1	3	\$37.19	\$12.40	1	11
Euthyrox TAB • DAW 0 = 142	257	\$1,058.28	\$4.12	142	1 – 64
 SYNTHROID (MSB) DAW 0 = 2 DAW 1 = 106 DAW 2 = 8 	393	\$18,360.88	\$46.72	116	0 – 63
TIROSINT CAP (SSB) • DAW 0 = 2 • DAW 1 = 2 • DAW 2 = 2	15	\$1,910.48	\$127.37	6	39 – 49
TIROSINT SOL (SSB) • DAW 0 = 6	21	\$3,000.38	\$142.88	6	1-41

^{*}Red font denotes brand utilization

Opioid Summary



- 1Q2018 to 4Q2019 excludes IHS
- 1Q2020 to current includes IHS
- March 13, 2020 Pandemic Closure

Opioid Initiatives:

- 1. June 1, 2018 early refill threshold for controlled substance changed from 75% to 85%
- 2. July 1, 2028 PA for more than one LAO and one SAO
- 3. August 1, 2018 opioid Naïve PA (initial 7-day supply and 60 MED limit)
- 4. October 1, 2018 to October 1, 2019 decrease from 300 MED to 90 MED (cancer diagnosis excluded)

Other Initiatives:

- Buprenorphine PA (Bunavail/Suboxone/Zubsolv/Subutex)/ST (Belbuca/Butrans) removed 10/14/2019
- Lidoderm PA removed 8/1/2020

Total Eligibility and Utilizers

TOTAL FIIGIDIII	ity and Othizers		
Quarter	Avg eligible	Avg utilizing members	% utilizing members of
Quarter	members	of all drugs	all drugs
1Q2020	123,573	27,089	21.9%
2Q2020	126,777	20,747	16.4%
3Q2020	132,373	23,417	17.7%
4Q2020	136,262	23,488	17.2%
1Q2021	139,748	24,405	17.5%
2Q2021	142,872	26,162	18.3%
3Q2021	146,023	27,847	19.1%
4Q2021	149,034	29,257	19.3%
1Q2022	151,735	28,892	19.0%
2Q2022	154,608	28,338	18.3%
3Q2022	157,627	29,109	18.5%

3Q2022

Jun 21 to Sep 21

Opioid Utilization Snapshot

Jun 22 to Sep 22

8

Opioid Claims 10,070

3.1% prescription claims filled for an opioid

0.5% higher than Medicaid FFS benchmark

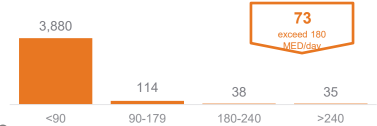


Utilizers **4,067 29.8%** are high utilizers

-4.5% lower than high utilizers Medicaid FFS

Utilizers by Cumulative MED4

Current CDC Guidelines⁵ urge doses of 90 MME⁶ or less in chronic opioid utilizers⁵





Shoppers: Poly Pharmacy

50 opioid utilizing members with 3+ pharmacies

Shoppers: Poly Prescriber

262 Shoppers: Poly Prescriber opioid utilizing members with 3+ prescribers



Opioid Claims 10,688

3.1% prescription claims filled for an opioid

0.9% higher than Medicaid FFS benchmark



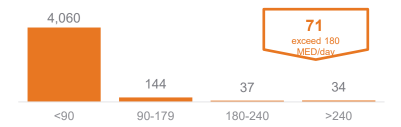
Utilizers 4,275

30.1% are high utilizers

0.5% higher than high utilizers Medicaid FFS

Utilizers by Cumulative MED⁴

Current CDC Guidelines⁵ urge doses of 90 MME⁶ or less in chronic opioid utilizers⁵





Shoppers: Poly Pharmacy

49 opioid utilizing members with 3+ pharmacies



Shoppers: Poly Prescriber

357 Shoppers: Poly Prescriber opioid utilizing members with 3+ prescribers



Opioid Utilization

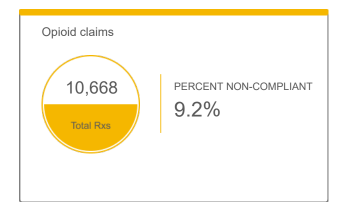
Opportunities date range: Jun - Sep 2022 Benchmark: MEDICAID FEE FOR SERVICE

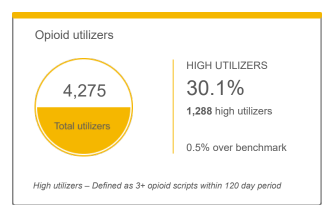
Utilizers: 4,275

3.1% of all Rx claims are filled for an Opioid

Opioid dependence can start in just a few days, and the risk of chronic opioid use increases with each additional day of opioid supplied, starting with the third day. Our Opioid Risk Management program, which includes point of sale, utilization management and retrospective drug utilization edits, are tightly aligned with CDC opioid prescribing guidelines which can help reduce exposure to excessive doses and prevent more members from transitioning from acute to chronic use.

- · Opioid prescriptions account for 3.1% of all prescriptions this period, which is 0.9% higher than the benchmark
- · 1,288 high opioid utilizers were identified this period, which is 0.5% higher than the benchmark





Claim breakdown



76.2% of all opioid Rxs were filled for short acting opioids. 1,863 Rxs were for medication assisted therapy (MAT) and 189 were for rescue therapy. CDC guidelines advise prescribers to manage pain with the lowest effective dose and to avoid or carefully justify doses for chronic users >90mg MME/day.

MAT – Medication Assisted Therapy (buprenorphine, etc) Overdose rescue therapy – opioid overdose reversals winaloxone MME – relative potency of an opioid to a morphine dose

Utilizers by cumulative MED

utilizers exceed 180 MED/day

MED Scores	<90	90-179	180-240	>240
Utilizers	4,060	144	37	34

MED – Morphine Equivalent Dose is a relative potency of an opioid to standard of a morphine; Cumulative MED is daily MED or narcotic load across all active opioid prescriptions in a members profile within a 120 day period

Language Assistance / Non-Discrimination Notice

Asistencia de Idiomas / Aviso de no Discriminación

語言協助 / 不歧視通知

Opioid Opportunity Assessment

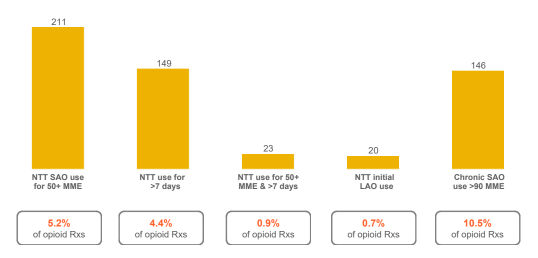
Opportunities date range: Jun - Sep 2022

Benchmark: MEDICAID FEE FOR SERVICE

Percent non-compliant: 9.2%

Utilizers non-compliant to opioid Rx CDC guidelines

(new to therapy and chronic use)



 $\textit{NTT-} \underbrace{\textit{view definition}} \ | \ \ \textit{SAO-} \underbrace{\textit{view definition}} \ | \ \ \textit{LAO-} \underbrace{\textit{view definition}} \ | \ \ \textit{MME-} \underbrace{\textit{view definition}}$



742

DID YOU KNOW?

49 opioid utilizing members use 3 or more pharmacies and 357 opioid utilizing members use 3 or more prescribers.

NNT – New to Therapy

SAO - Short Acting Opioid

LAO - Long Acting Opioid

MME – Morphine Milligram Equivalent represents a relative potency of an opioid to a morphine dose

Opioid utilizers with potentially contraindicated medication use

SKELETAL MUSCLE RELAXANTS BENZODIAZEPINES ANTICONVULSANTS MEDICATION ASSISTED THERAPY PRENATAL

766

Anticonvulsants – gabapentin, pregabalin, Anticonvulsant benzodiazepines (clobazam, clonazepam, diazepam)

551

Language Assistance / Non-Discrimination Notice

Asistencia de Idiomas / Aviso de no Discriminación

語言協助 / 不歧視通知

122

N/A

Table of Contents

1.	Administrative PA	
	a. DAW	24
	b. General PA	25
	c. Max Units Override	26
	d. Medications > \$5000	
2.	Acne Agents Topical	
3.	Rosacea Agents Topical	
4.	Allergen Extract (Grastek, Oralair, Ragwitek)	
5.	Altabax	
6.	Antidepressants	32
7.	Brisdelle	33
8.	Antipsychotics	34
9.	Antiemetics: Akynzeo/ Bonjesta/Diclegis/ Sancuso/ Zuplenz	35-39
10.	Antihistamines: Non-Sedating Antihistamines	40
	. ARBS (Edarbi, Edarbyclor)	
12.	. Amrix, Fexmid	42
13.	Brexafemme	43
14.	. Cambia, Zipsor, Zorvolex	44
15.	Chronic Constipation Agents (Amitiza, Linzess, Movantik)	45
16.	CGRP Inhibitors (Aimovig, Ajovy, Emgality)	46-47
17.	Desoxyn	48
18.	Dificid	49
19.	Durlaza	50
20.	. Emflaza	51
21.	Epidiolex	52
22.	Evrysdi	53-55
23.	Genitourinary Smooth Muscle Relaxant	56
24.	GLP-1 Agonists	57
25.	Gralise, Horizant	58
26.	Growth Hormone: Adult-Pediatric, Serostim, Zorbtive	59-63
27.	. Head Lice Medication	64
28.	. Hemangeol	65
29.	. Hepatitis C	66-67
30.	Brand Name Narcotics	68-69
31.	. Hydrocodone-APAP	70-71
32.	Opioid MED Limit	72
	. Opioid Naïve	
	. Opioid LAO/SAO	
	· Idiopathic Pulmonary Fibrosis (Esbriet, Ofev)	
	TIM, JAK, Monoclonal Antibody, etc	
	a. Actemra	76-77
	b. Adbry	
	c. Cibingo	
	d. Cimzia	
	e. Cosentyx	82
	f. Dupixent	
	g. Enbrel	
	h. Enspryng	
	i. Fasrena	
	j. Humira	
	k. Ilaris	
	l. Ilumya	
	m. Kevzara	
	n Vinoret	

	o. Nucala	95
	p. Olumiant	96
	q. Orencia	97-98
	r. Otezla	99
	s. Rinvoq	100
	t. Siliq	101
	u. Simponi	
	v. Skyrizi	
	w. Sterlara	
	x. Taltz	
	y. Tremfya	
	z. Xeljanz/XR	
	aa. Xolair	
27	Juxtapid	
	Ketoconazole Agents Topical (Extina, Xolegel/Duo)	
	Onychomycosis Agents Topical Luzu	
	Oravig	
	Vusion	
	Makena	
	Metozolv	
	Multiple Sclerosis	
	Tysabri	
	Nasal Steroids	
	Nascobal	
	Nuplazid	
50.	Nuvessa	125
51.	Hetlioz	126
52.	Nuvigil, Provigil	127
53.	Sunosi, Wakix	128
54.	Xyrem	129
55.	Onfi	130
56.	Ophthalmic Antihistamines (Bepreve, Lastacaft, Pataday, Patanol, Pazeo)	131
57.	Opzelura	132
58.	Oracea, Solodyn, Seysara	133
	Otrexup	
60.	PCSK9 Inhibitors (Praluent, Repatha)	135
	Proton Pump Inhibitors	
	Duexis, Vimovo	
	Qualaquin	
	Rayos	
	Relistor	
	Soma 250	
	Tivorbex	
	Tramadol: Ultram ER, tramadol ER/SR, Conzip, Synapryn	
	Triptans	
	Onzetra/Xsail	
	Nurtec ODT, Reyvow, Ubrelvy	
	Uloric	
	Viberzi	
	Xenazine	
	Xepi	
	Xifaxan	
	Zolpidem (Ambien CR, Edluar, Intermezzo SL, Zolpimist)	
78.	Belsomra, Dayvigo, Quiviq	154



Dispense As Written (DAW) Prior Authorization Request Form

Member Information (required)			Pr	Provider Information (required)			
Member Name:			Provider Nam	Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:		l l		
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street	Address:			
Phone:	"	<u> </u>	City:	State:		Zip:	
		Medication I	Information (re	equired)			
Medication Name:			Strength:		Dosage F	orm:	
•	Check if requesting brand		Directions for	Use:			
□ Check if request	is for continuatio	n of therapy					
		Clinical Inf	formation (requ	uired)			
Clinical inform	ation:						
•		failure with the generi	•				
Has the patient must be comple		the generic product a ☐ No	nd experienced a	n adverse reac	tion (a Med	Watch form	
Does the patien	nt have a contra	indication to the gene	ric product? 🗖 Y	es 🗆 No			
Is the generic p	roduct unavaila	ble? 🛚 Yes 🖵 No					
Are there any other to this review?	comments, diagnos	es, symptoms, medications	tried or failed, and/or	any other information	on the physici	an feels is important	
Please note: T	his request may be o	denied unless all required info	rmation is received.				

For urgent or expedited requests please call 1-855-401-4262.



Please note: All information below is required to process this request.

Fax to 1-844-403-1029

Mon-Sat: 7am to 7pm Central

Prior Authorization Request Form

		FOR FUTURE USE. FORMS					
Memb	er Inform	nation (required)	Pro	Provider Information (required)			
Member Name:			Provider Name	:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Ad	ddress:			
Phone:	<u> </u>	I	City:	State:	Zip:		
		Medication	Information (red	quired)			
Medication Name:		Mediedilen	Strength:	quireu)	Dosage Form:		
☐ Check if requesting	brand		Directions for U	Jse:			
☐ Check if request is		n of therapy	23000710707				
		Clinical In	formation (requir	red)			
What is the patient	t's diagnosis	for the medication bei	ing requested?				
			ICD-10 Code	a(e):			
What medication(s	the nat	ient tried and failed?	100-10 0000	5(3)			
Are there any supp	oorting labs o	or test results? (Please	e specify)				
☐ Titration or loadi☐ Patient is on a d bedtime)☐ Requested stren	requested per a for exceeding ng dose purpo ose-alternating gth/dose is no	ng the plan limitations′ oses	blet in the morning ar		night, one to two tablets at		
Are there any other corto this review?	mments, diagnos	ses, symptoms, medications	s tried or failed, and/or a	ny other informatio	n the physician feels is important		
Please note: This	request may be	denied unless all required info	ormation is received.				

For urgent or expedited requests please call 1-855-401-4262.



Fax to 1-844-403-1029

Mon-Sat: 7am to 7pm Central

Quantity Limit Request Form

Memb	er Informatio	TURE USE. FORMS ARE (der Infori		
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Addres	SS:		
Phone:			City:	State:		Zip:
		Medication Info	rmation (required	i)		
Medication Name:			Strength:	·	Dosage Fo	orm:
☐ Check if requesting			Directions for Use:		<u> </u>	
☐ Check if request is	for continuation of th					
		Clinical Inforr	nation (required)			
What is the patient	's diagnosis for the	e medication being re	equested?			
			ICD-10 Code(s): _			
What is the quantity	requested per DAY	?				
bedtime) ☐ Requested stren	ng dose purposes ose-alternating sche gth/dose is not comi a greater quantity fo	dule (e.g., one tablet in mercially available r the treatment of a lar	-			
Are there any other cor to this review?	nments, diagnoses, syr	nptoms, medications tried	or failed, and/or any ot	her information	n the physicia	an feels is important
Please note: This	request may be denied u	nless all required informatio	n is received.			

For urgent or expedited requests please call 1-855-401-4262.



High Dollar/Claim Dollar Amount Override Prior Authorization Request Form

Member Information (required)				Provider Information (required)			
Member Name	e:		Provider Nam	e:			
Insurance ID#	:		NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address	S:		Office Fax:				
City:	State:	Zip:	Office Street A	Address:			
Phone:			City:	State:	Zip:		
		Medication	Information (re	equired)			
Medication Na	ime:		Strength:	. ,	Dosage Form:		
☐ Check if red	questing brand		Directions for	Use:	1		
☐ Check if red	quest is for continuatio r	of therapy					
		Clinical In	formation				
			formation (requ	•			
What is the	patient's diagnos	is for the medicatio	on being requeste	ed?			
			ICD-10 Code	(s):			
What is the	requested quanti	ty per day/fill/presc	ription/ or month	?			
					onth and the duration		
(i.e., 3 caps information.		sules per prescriptior	n/per 30 days). Use	e/take as directe	d is not sufficient		
Are there any of to this review?	other comments, diagnos	es, symptoms, medications	s tried or failed, and/or	any other informatior	n the physician feels is importan		
Please note:		enied unless all required info					



Topical Acne Agents Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)				Provider Information (required)			
Member Name:			Provider Na	me:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phon	e:			
Street Address:			Office Fax:				
City:	State:	Zip:	Office Stree	t Address:			
Phone:			City:	State:	Zip:		
		Medication	Information	(required)			
Medication Name:			Strength:	(Dosage Form:		
☐ Check if requesting	ng brand		Directions fo	or Use:			
☐ Check if request in	s for continuation	on of therapy					
		Clinical In	formation (required)			
Select the diagno	sis below:						
☐ Acne vulgaris							
Plaque psorias	is [Tazorac (ta	zarotene) only]					
Other diagnosis	s:		IC	D-10 Code(s):			
Medication histor	ry:						
		lure of a generic topical a um/sulfur, sulfacetamide			noin, clindamycin phosphate, ′es □ No		
Are there any other conthis review?	mments, diagnos	es, symptoms, medications t	ried or failed, and/o	or any other information	on the physician feels is important to		
Please note: This	request may be d	enied unless all required inform	nation is received.				

For urgent or expedited requests please call 1-855-401-4262.



Topical Rosacea Agents Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address	3:		
Phone:	l		City:	State:		Zip:
		Medication Info	ormation (require	ed)		
Medication Name:			Strength:		Dosage Fo	orm:
☐ Check if requesting			Directions for Use:			
Check if request is f	for continuation of the	ару				
		Clinical Infor	mation (required)			
Select the diagnos	is below:					
Acne rosacea						
□ Other diagnosis:	-		ICD-10 Cod	de(s):		
Medication history	:					
Has the patient had	a trial of a generic top	oical acne agent (benz	zoyl peroxide, clinda	mycin phosp	hate, eryth	romycin,
		de sodium, tretinoin, m	netronidazole cream	/gel/lotion) ir	the past	
120 days? ☐ Yes [⊒ No					
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?						
			····			
Please note: This	request may be denied un	ess all required information	n is received.			

For urgent or expedited requests please call 1-855-401-4262.



Grastek®, Oralair®, Ragwitek® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)		Provider Information (required)			
Member Name:			Provider Name	:	
Insurance ID#:			NPI#: Specialty:		
Date of Birth:			Office Phone:		
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street Ad	ddress:	
Phone:	1		City:	State:	Zip:
		Medication I	nformation (re	equired)	
Medication Name:			Strength:	,,,,,,	Dosage Form:
☐ Check if requesting	brand		Directions for U	Jse:	
☐ Check if request is	for continuation of t	herapy			
		Clinical Info	ormation (requ	ıired)	
What is the patient	t's diagnosis for th	ne medication bein			
				,	
ICD-10 Code(s): _					
Clinical informatio	n:				
Is the patient's diag	nosis confirmed by	a positive skin test o	or in vitro testing fo	r pollen-specific	IgE antibodies? ☐ Yes ☐
Has the patient had No	a history of failure	or intolerance to sub	cutaneous allerge	n immunotherap	y (allergy shots)? 🗖 Yes 🗖
Does the patient ha	ve severe, unstable	e or uncontrolled astl	nma? 🗆 Yes 🗅 N	lo	
Select the medicat	ion categories tha	nt the patient has tr	ied and failed:		
	, -	astine, olopatadine,		•	
☐ Intranasal cortico triamcinolone)	osteroids (e.g., bec	lomethasone, budes	onide, ciclesonide	, flunisolide, flutio	casone, mometasone,
☐ Leukotriene inhib	bitors (e.g., montelu	ıkast, zafirlukast, zile	euton)		
Oral antihistaming	nes (e.g., cetirizine,	desloratadine, fexof	enadine, levocetiri	izine, or loratadir	ne)
Are there any other com this review?	ments, diagnoses, syn	nptoms, medications tri	ed or failed, and/or ar	ny other information	n the physician feels is important to
Please note: This r	equest may be denied u	nless all required informa	tion is received.		

For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.



Altabax® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)					
Member Name:			Provider Name:					
Insurance ID#:			NPI#:	NPI#: Specialty:				
Date of Birth:			Office Phone:					
Street Address:			Office Fax:	Office Fax:				
City:	State:	Zip:	Office Street Address:					
Phone:	none:		City:	State:	Zip:			
		Medication In	formation (re	equired)				
Medication Name:			Strength:	<i>-</i>	Dosage Form:			
☐ Check if requesting	brand		Directions for U	Jse:				
☐ Check if request is	for continuation of t	herapy						
		Clinical Info	rmation (requ	ıired)				
Select the diagno	sis below:							
_		cus aureus (MRSA)						
Other diagnosi	s:		ICD-10 Code(s):					
Medication histo	ry:							
Has the patient trid days? ☐ Yes ☐		eric mupirocin ointm	ent or cream fo	or a minimum o	of 5 days within the last 90			
Quantity limit red	•	MACNITUO						
What is the quanti	•							
		, the plan limitatio r y to cover a larger s						
Other:	o a largor quartit	y to oover a larger of	arrado arca					
Are there any other community this review?	ments, diagnoses, syr	nptoms, medications tried	l or failed, and/or ar	ny other informatio	on the physician feels is important to			
Please note: This re	nauget may be desired.	unless all required information	on is received					

For urgent or expedited requests please call 1-855-401-4262.



Antidepressants Prior Authorization Request Form

Member Information (required)				Provider Information (required)		
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Add	Office Street Address:		
Phone:	I	I	City:	State:	Zip:	
		Medication	on Information (req	uired)		
Medication Name	e:		Strength:	,	Dosage Form:	
☐ Check if reque	esting brand		Directions for Use	e:		
☐ Check if reque	est is for continuatio	n of therapy				
		Clinical	Information (require	ed)		
What is the pat	tient's diagnosis fo	the medication beir	ng requested?			
		ICI	D-10 Code(s):			
Clinical inform	nation:					
Is the patient a	Iready stabilized o	n therapy with the re	quested medication? Y	'es □ No		
Please list ALL	medications the p	atient has had a trial	of within the past 12 mor	nths:		
concentrate re	equests, also ans	wer the following:	spension, Prozac solution		olTab, and Zoloft	
Quantity limit						
•		er DAY?				
	ason for exceed!! loading dose purpo	ng the plan limitatio	ons?			
□ Patient is or			e tablet in the morning and	d two tablets at	night, one to two tablets at	
bedtime)	otropath/doop is no	ot commorcially avail	lahla			
Other:	strength/dose is no	ot commercially avail				
Are there any other	r comments, diagnos	es, symptoms, medication	ons tried or failed, and/or any	other information	the physician feels is important t	
Please note:	This request may be d	enied unless all required i	nformation is received.			

For urgent or expedited requests please call 1-855-401-4262.



Brisdelle™ Prior Authorization Request Form

Member Information (required)			Pı	Provider Information (required)				
Member Name:			Provider Name	Provider Name:				
Insurance ID#:			NPI#:	NPI#: Specialty:				
Date of Birth:			Office Phone:	Office Phone:				
Street Address	S:		Office Fax:	Office Fax:				
City:	City: State: Zip:			Office Street Address:				
Phone:	I	L	City:	State:	Zip:			
		Medication	Information (required)				
Medication Name:			Strength:	Dosage Form:				
☐ Check if requesting brand			Directions for	Directions for Use:				
☐ Check if req	uest is for continuatio	n of therapy						
		Clinical In	formation (red	uired)				
Medication	history:							
Has the pati	ient had a 60 day tı	rial and failure of parc	exetine oral tablets	s within the past	6 months? U Yes U No			
Are there any oth	ner comments, diagnose	es, symptoms, medications	tried or failed, and/or a	any other information	n the physician feels is important to			
Please note:	This request may be de	enied unless all required infor	mation is received.					

For urgent or expedited requests please call 1-855-401-4262.



Atypical Antipsychotics Prior Authorization Request Form

Member Information (required)			Pro	Provider Information (required)			
Member Name:			Provider Name:	Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Add	Office Street Address:			
Phone:			City:	State:	Zip:		
		Medicatio	n Information (requ	uired)			
Medication Nam	ne:		Strength:	·	Dosage Form:		
☐ Check if requ	esting brand		Directions for Use	e:			
Check if requ	est is for continuatio r	of therapy					
		Clinical	Information (require	ed)			
clinical inform or patients with or patients you eurologist invo	h a diagnosis of depresunger than 6 years of a blved in care? Yes	ssion, has the patient tr ge, is a psychiatrist, de No	ried and failed 2 different ar evelopmental pediatrician, c , injectables, extended-re	:hild/adolescen	t psychiatrist or pediatric		
•	nable to swallow?		class in the last 30 days?	J Voo □ No			
Quantity limit r What is the qua What is the rea Titration or lo Patient is on	requests: Intity requested per DA ason for exceeding the Dading dose purposes	Y?e plan limitations?			ne to two tablets at bedtime)		
	er comments, diagnoses	s, symptoms, medication	ns tried or failed, and/or any	other information	on the physician feels is importan		
lease note:	This request may be de	nied unless all required in	formation is received.				

For urgent or expedited requests please call 1-855-401-4262.



Akynzeo® Prior Authorization Request Form

Member Information (required)			Pr	Provider Information (required)				
Member Name:			Provider Name:					
Insurance ID#:			NPI#:		Specialty:			
Date of Birth:			Office Phone:					
Street Address	Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Ac	Office Street Address:				
Phone:	I	I	City:	State:	Zip:			
		Medication	Information (r	equired)				
Medication Na	nme:		Strength:		Dosage Form:			
☐ Check if red	questing brand		Directions for U	Directions for Use:				
☐ Check if red	quest is for continuatio	n of therapy						
		Clinical In	nformation (requ	uired)				
Select the	diagnosis below:							
☐ Prophyla	axis of chemothera	oy-induced nausea/v	omiting					
☐ Other diagnosis:			ICD-10 Co	ode(s):				
Clinical inf	ormation:							
		emetogenic chemot 90 days? ☐ Yes ☐		or regimens inc	cluding anthracyclines and			
Are there any oth	her comments, diagnose	es, symptoms, medications	s tried or failed, and/or a	ny other information	on the physician feels is important to			
Please note:	This request may be de	enied unless all required info	rmation is received.					

For urgent or expedited requests please call 1-855-401-4262.



Bonjesta® Prior Authorization Request Form

Member Information (required)				Provider Information (required)			
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address	:		Office Fax:				
City:	State:	Zip:	Office Street A	Office Street Address:			
Phone:			City:	State:	Zip:		
		Medication	Information (required)			
Medication Nar	me:		Strength:		Dosage Form:		
☐ Check if requ	uesting brand		Directions for	Use:			
☐ Check if requ	uest is for continuatio	n of therapy					
		Clinical In	nformation (req	uired)			
☐ Hypereme ☐ Other diag Quantity limi What is the qu What is the r ☐ Titration or ☐ Patient is or tablets at b ☐ Requested	it requests: uantity requested pereason for exceeding lose purpoon a dose-alternating dedtime) d strength/dose is no	ng the plan limitations	? ablet in the morning				
Are there any oth this review?	er comments, diagnose	es, symptoms, medications	tried or failed, and/or a	any other informatio	n the physician feels is important to		
Please note:		enied unless all required infor					



Diclegis® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)				
Member Name:			Provider Nam	e:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street A	Address:			
Phone:			City:	State:	Zip:		
		Medication Inf	ormation ((required)			
Medication Name:			Strength: Dosage Form:				
☐ Check if requestin	•		Directions for Use:				
☐ Check if request is	for continuatio	n of therapy					
		Clinical Info	rmation (red	quired)			
Select the diagn	osis below:						
Hyperemesis	gravidarum						
Other diagnos	sis:		ICD-10 Co	ode(s):			
Are there any other co	Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to						
		denied unless all required information					

For urgent or expedited requests please call 1-855-401-4262.



Sancuso® Prior Authorization Request Form

	DO NOT COPY FOR FU	TURE USE. FORMS ARE U	JPDATED FREQUENTLY	AND MAY BE	BARCODED
Memb	er Informatio	N (required)	Provid	der Info	rmation (required)
Member Name:			Provider Name:		
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone:		
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street Address:		
Phone:		1	City:	State:	Zip:
		Medication Inf	ormation (required	d)	
Medication Name:			Strength:	<u>′</u>	Dosage Form:
☐ Check if requesting	brand		Directions for Use:		
☐ Check if request is f	for continuation of th	erapy			
		Clinical Infor	mation (required)		
Select the diagnos	is helow:		(,)		
☐ Prophylaxis of ch		d nausea/vomiting			
	• •	•	ICD-10 Code(s):		
Clinical information			. ,		
Has the patient had days? ☐ Yes ☐ No		Hydroxytryptamine type	e 3 (5-HT3) receptor a	antagonist f	or 14 days in the past 90
Is the patient receiving days? ☐ Yes ☐ No.	•	or highly emetogenic cl	nemotherapy for up to	5 consecu	tive
Is the patient unable difficulty swallowing		dications for chemother	rapy-induced nausea	and vomitin	ng due to a diagnosis of
Quantity limit requ					
What is the quantity	•				
What is the reason		plan limitations?			
☐ Titration or loadin☐ Patient is on a do		dule (e.g., one tablet ir	the morning and two	tablets at r	night, one to two
tablets at bedtime	e)	. •			,
Requested stren	gth/dose is not comi	mercially available			
Other:					
Are there any other conthis review?	nments, diagnoses, syn	nptoms, medications tried	or failed, and/or any othe	er information	n the physician feels is important to
Discourantes This					

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262.



Zuplenz® Prior Authorization Request Form

Member Information (required)			Pı	Provider Information (required)			
Member Name	:		Provider Name	:			
Insurance ID#:			NPI#: Special				
Date of Birth:			Office Phone:				
Street Address:	:		Office Fax:	Office Fax:			
City:	State:	Zip:	Office Street A	ddress:			
Phone:			City:	State:	Zip:		
		Medication	n Information (r	equired)	·		
Medication Nar	ne:		Strength:		Dosage Form:		
☐ Check if requ	uesting brand		Directions for U	Jse:			
☐ Check if requ	uest is for continuatio r	of therapy					
		Clinical I	nformation (req	uired)			
Clinical info	ormation:						
•	ent had a trial of a q	generic -Hydroxytryp	tamine type 3 (5-H	T3) receptor an	tagonist for 14 days in the		
Is the patien	t receiving moderat	tely and/or highly em	netogenic chemothe	erapy for up to 5	consecutive		
days? 🛚 Ye	es 🗆 No						
Are there any ot this review?	ther comments, diagnos	es, symptoms, medication	s tried or failed, and/or a	ny other informatio	n the physician feels is important to		
Please note:	This request may be d	enied unless all required info	ormation is received.				

For urgent or expedited requests please call 1-855-401-4262.



Non-Sedating Antihistamines Prior Authorization Request Form

Member Information (required)			Pi	Provider Information (required)			
Member Name:			Provider Name	Provider Name:			
Insurance ID#:			NPI#:	NPI#: Specialty:			
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street A	Office Street Address:			
Phone:			City:	State:	Zip:		
		Medication	Information	(required)			
Medication Name	y:	Medication	Strength:	required)	Dosage Form:		
☐ Check if reque	sting brand		Directions for	Use:			
	st is for continuatio	n of therapy					
		Clinical I	nformation (red	auired)			
Medication his Has the patient fexofenadine & Clinical inform	osis: tory: tried and failed a 1 pseudoephedrine, ation:	loratadine, or loratadi	ne following: cetirizin ne & pseudoephedri	e, cetirizine & pse ine? □ Yes □ N	eudoephedrine, fexofenadine,		
		ted difficulty in swallov	ving diagnosis? 🔲 🕽	res U No			
What is the rea ☐ Titration or lo ☐ Patient is on bedtime) ☐ Requested s ☐ Other:	ntity requested penson for exceeding dose purpo a dose-alternating strength/dose is no	g the plan limitations ses g schedule (e.g., one to t commercially availab	ablet in the morning		at night, one to two tablets at		
Are there any other his review?	comments, diagnose	s, symptoms, medications	s tried or failed, and/or a	any other informatio	n the physician feels is important to		
Please note:	This request may be de	nied unless all required info	rmotion is resolved				

For urgent or expedited requests please call 1-855-401-4262.



Edarbi and Edarbyclor Prior Authorization Request Form

Me	mber Inform	ation (required)			ermation (required)
Member Name:			Provider Nan	ne:	
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone	:	
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street	Address:	
Phone:			City:	State:	Zip:
		Medication	Information	(required)	
Medication Name	э:		Strength:	<u> </u>	Dosage Form:
☐ Check if reque	esting brand		Directions for	· Use:	
☐ Check if reque	est is for continuatio	n of therapy			
		Clinical Ir	nformation (re	equired)	
Clinical infor	mation:				
Has the patier days?		the requested angio	otensin II recep	otor blocker (A	RB) for more than 60
Has the patier days?	•	ensin-converting enzy	yme (ACE) inhibi	tor or a generic	ARB within the last 120
	ent have an additrenal failure?	tional diagnosis of chi	ronic obstructive	pulmonary dise	ase (COPD) or
Are there any other this review?	comments, diagnose	es, symptoms, medications	tried or failed, and/or	any other information	on the physician feels is important to
		enied unless all required infor			



Amrix® & Fexmid® (cyclobenzaprine) Prior Authorization Request Form

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:		1	
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Add	Office Street Address:		
Phone:		<u>I</u>	City:	State:	2	Zip:
		Medication	Information (red	guired)		
Medication Name:			Strength:	-{·····	Dosage For	m:
☐ Check if requesting	brand		Directions for Us	se:		
☐ Check if request is	for continuation (of therapy				
		Clinical In	formation (requi	red)		
Select the diagno	osis below:		` '	,		
_		herapy for relief of r	nuscle spasm asso	ciated with ac	ute. painful	musculoskeletal
conditions	and priyologic		naccio opacini acce	olatoa Willia	ato, pannar	accarconora
	s:		ICD-10 Cod	de(s):		
Medication histo						
	•	dav trial and failure	e of cyclobenzaprine	e 5 mg tablets	OR cyclobe	enzaprine 10
		ays? 🗆 Yes 🗅 No				
Quantity limit rec	quests:					
		er DAY?				
		ing the plan limitat	tions?			
☐ Titration or load						
		ng schedule (e.g., o	ne tablet in the mo	rning and two	tablets at ni	ght, one to two
tablets at bedti	,	ot commercially av	ailahla			
Other:	-	-	allabic			
Are there any other comi his review?	ments, diagnoses,	symptoms, medications	tried or failed, and/or any	y other information	n the physician	feels is important to
						
Please note: This re	equest may be deni	ed unless all required infor	mation is received			

For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.



Brexafemme® Prior Authorization Request Form

		R FUTURE USE. FORMS A				
Member Name:	er Informa	(ICII (required)	Provider Information (required) Provider Name:			
				•		
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Ac	ddress:		
Phone:			City:	State:	Zip:	
		Medication	Information (re	equired)		
Medication Name:			Strength:		Dosage Form:	
☐ Check if requesting	g brand		Directions for U	Jse:		
☐ Check if request is		of therapy				
		Clinical In	formation (requ	uired)		
Select the diagnos	sis below:		(,		
☐ Vulvovaginal ca						
☐ Other diagnosis			_ ICD-10 Code(s): _			
Clinical information	on:					
Has the patient trie	d and failed 3 tri	als of fluconazole or cl	otrimazole in the pa	ıst 14 days? 🗖 \	∕es □ No	
Quantity limit requ What is the quantity		MONTH?				
		the plan limitations?	?			
☐ Titration or load				4 4	. 4 minute - m - 4 - 4 - m	
tablets at bedtim		schedule (e.g., one tab	plet in the morning a	ind two tablets a	at night, one to two	
		commercially available)			
Other:	-	•				
Are there any other com this review?	nments, diagnoses	, symptoms, medications t	ried or failed, and/or ar	ny other informatio	n the physician feels is important to	

This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262. Please note:



Cambia[®], Zipsor[®], Zorvolex[®] Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required) Provider Information (required) Member Name: Provider Name: NPI#: Insurance ID#: Specialty: Office Phone: Date of Birth: Street Address: Office Fax: City: Office Street Address: State: Zip: Phone: City: State: Zip: Medication Information (required) Strength: Medication Name: Dosage Form: ☐ Check if requesting brand Directions for Use: ☐ Check if request is for continuation of therapy Clinical Information (required) **Medication history:** Has the patient had a documented 30 day trial of a generic diclofenac product within the last 120 days? ☐ Yes ☐ No Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?

Please note:

This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Amitiza®, Linzess®, MovantikTM Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)		Provider Information (required)				
Member Name:		Provider Name:				
Insurance ID#:			NPI#: Specialty:			
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:	l	1	City:	State:		Zip:
		Medication Inf	ormation (required)		
Medication Name:			Strength:		Dosage Fo	orm:
☐ Check if requesting			Directions for Use:			
☐ Check if request is t	for continuation of the					
		Clinical Infor	mation (required)			
☐ Irritable bowel ☐ Opioid-induced ☐ Other diagnosis For opioid-induced Is the pain associate Quantity limit recommendation	thic constipation [Aisyndrome with constipation in an action of the constipation in an ated with cancer? [Aisyndrome constipation in ated with cancer? [Aisyndrome constipation]	adult patient with ch an adult patient w ☑ Yes ☑ No	only] mitiza and Linzess on the pain [Amitiza and Linzes on the pain [Amitiza and Code(s): _ ith chronic pain, and the pain and the pai	and Mov a		
☐ Titration or load ☐ Patient is on a tablets at bedtii ☐ Requested street	What is the quantity requested per DAY? What is the reason for exceeding the plan limitations? □ Titration or loading dose purposes □ Patient is on a dose-alternating schedule (e.g., one tablet in the morning and two tablets at night, one to two tablets at bedtime) □ Requested strength/dose is not commercially available □ Other:					
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is impthis review?				an feels is important to		

<u>Please note:</u> This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Aimovig[™], Ajovy[™], Emgality[™] Prior Authorization Request Form (Page 1 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)				
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Address:				
Phone:			City:	State:		Zip:	
		Medication Inf	ormation (required	t)			
Medication Name:			Strength:	-,	Dosage F	orm:	
☐ Check if requesting	brand		Directions for Use:				
	for continuation of the	erapy					
		Clinical Infor	mation (required)				
Select the diagnosis	below:						
☐ Chronic migraines							
☐ Episodic migraines	3						
Other diagnosis: _			ICD-10 Cod	de(s):			
Clinical information:							
Is the requested medi	cation prescribed by or	in consultation with a n	eurologist or pain/heada	ache specia	list? Yes	i □ No	
Will the requested me	dication be used in com	nbination with another C	GRP inhibitor? Yes	□ No			
	c therapies the patient holerance/contraindication		re, (defined as at least 2	? months of	therapy with	greater than 80%	
☐ Antidepressants (i.	.e., venlafaxine or tricyc	lic antidepressant such	as amitriptyline or nortri	iptyline)			
Please specify:							
☐ Anti-epileptics (i.e.	, topiramate or divalpro	ex sodium). Please spe	ecify:				
☐ Beta-blockers (i.e.	, atenolol, propranolol, r	nadolol, timolol, or meto	prolol). Please specify:				
For chronic migraine	es, also answer the fo	llowing:					
	evaluated for rebound h ISAIDs)? 🏻 Yes 🗘 No	neadaches caused by m	nedication overuse (more	e than 12 do	oses per mo	onth of narcotics,	
If diagnosed, will treat	ment include a plan to t	taper off the offending n	nedication? 🛚 Yes 🗆 N	lo			
Does the patient have months? Yes		o 15 headache days pe	r month, of which at lea	st 8 must be	e migraine d	lays for at least 3	
For episodic migrain	nes, also answer the fo	ollowing:					
Does the patient have 4 to 14 migraines per month (but no more than 14 headache days per month)? Yes No)		
Reauthorization:							
	ation request, answer						
Has the patient experi intensity? • Yes • N		nse to therapy, demons	trated by a reduction in	headache fi	requency ar	nd/or	
Has the use of acute i	migraine medications (e	e.g., NSAIDs, triptans, n	arcotics) decreased sind	ce the start	of CGRP th	erapy? 🗆 Yes 🗅 No	
Is the requested medication prescribed by or in consultation with a neurologist or pain/headache specialist? Yes No				i □ No			



Aimovig[™], Ajovy[™], Emgality[™] Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?									
Please note:	This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.								



Desoxyn® (methamphetamine) Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)		Provider Information (required)			
Member Name:			Provider Name	e:	
Insurance ID#:			NPI#: Specialty:		
Date of Birth:			Office Phone:		
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street Address:		
Phone:			City:	State:	Zip:
		Medication Inf	ormation (required)	
Medication Name:			Strength:		Dosage Form:
☐ Check if requesting	ng brand		Directions for	Use:	
☐ Check if request	is for continuation of th	erapy			
		Clinical Info	rmation (red	quired)	
Select the diagno	osis below:				
□ Attention Defice	it Disorder with Hypera	activity			
Other diagnosi	s:	1(CD-10 Code(s)	:	
medications from	ad a trial and failure (af any of the following op etine	otions in the past 90 dats ts product			tolerance to any four
Are there any other of this review?	comments, diagnoses, syn	nptoms, medications tried	or failed, and/or a	any other informatior	n the physician feels is important to
	nis request may be denied u or urgent or expedited reque				

This form may be used for non-urgent requests and faxed to 1-844-403-1029.

48



Dificid® Prior Authorization Request Form

Meml	ber Informa	OR FUTURE USE. FORMS A NTION (required)			EBARCODED Prmation (required)
Member Name:		rerorr (required)	Provider Nam		Titiation (required)
Insurance ID#:			NPI#:	Specialty:	
Date of Birth:			Office Phone	:	
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street	Address:	
Phone:			City:	State:	Zip:
		Medication	Information	(required)	
Medication Name:			Strength:	(roquirou)	Dosage Form:
☐ Check if requesting	g brand		Directions for	· Use:	
☐ Check if request is	for continuation	of therapy			
		Clinical Ir	nformation (re	equired)	
Select the diagn	osis below:				
□ Clostridium dif	fficile-associate	ed diarrhea (CDAD)			
Other diagnos	is:		ICD-10 Cod	le(s):	
Clinical informat	_				
•	•	the current guideline	es? 🛚 Yes 🗖 N	lo	
	•	atient has failed:			
•	•	ate severity) – metro	nidazole		
•	(severe) – van	•	n and matronida	7010	
•	•	licated) – vancomyci nen as first episode	n and metronida.	zoie	
	•	ncomycin in tapered	rogimon		
- Second recuir	erice – orai vai	iconiyeni in tapered	regimen		
Are there any other co this review?	mments, diagnose	s, symptoms, medications	tried or failed, and/or	any other informatio	n the physician feels is important to
Please note: This	e request may be do	nied unless all required infor	mation is received		

This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262.



DurlazaTM Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)				Provider Information (required)			
Member Name:			Provider Name	Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Ac				
Phone:			City:	City: State: Zip:			
		Medication	n Information (r	equired)			
Medication Name:			Strength:	· · · · · · · · · · · · · · · · · · ·	Dosage Form:		
☐ Check if requesting	brand		Directions for U	Jse:			
☐ Check if request is	for continuation	of therapy					
		Clinical I	nformation (requ	uired)			
Select the diagno	osis below:						
☐ Chronic corona	ary artery disea	ase (CAD)					
□ Ischemic strok	е						
☐ Transient ische	emic attack						
Other diagnosi	is:		ICD-10 Cod	de(s):			
Clinical informat	ion:						
Has the patient ha	ad a 90 day tria	al and failure with im	mediate release as	spirin? 🗖 Yes 🛭	⊒ No		
Please submit clin	nical rationale e	explaining why a fail	ure with the extend	ed-release prod	duct is not expected:		
Are there any other corthis review?	mments, diagnose	s, symptoms, medication	s tried or failed, and/or a	ny other information	n the physician feels is important to		
Please note: This		nied unless all required info					

For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.



Emflaza[™] Prior Authorization Request Form

		OR FUTURE USE. FURMS A						
Mem	ber Informa	ntion (required)	P	Provider Information (required)				
Member Name:			Provider Nam	Provider Name:				
Insurance ID#:			NPI#:		Specialty:			
Date of Birth:			Office Phone:					
Street Address:			Office Fax:					
City: State: Zip:			Office Street	Address:				
Phone:	'	<u> </u>	City:	Zip:				
		Medication	Information	(required)				
Medication Name:			Strength:	ngth: Dosage Form:				
☐ Check if requestir	ng brand		Directions for	Directions for Use:				
☐ Check if request i	s for continuation	of therapy						
		Clinical In	formation (re	quired)				
Select the diagr	nosis below:							
□ Duchenne mu	uscular dystroph	ny						
Other diagnos	sis:		ICD-10 Cd	ICD-10 Code(s):				
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?								
Fo	r urgent or expedited	nied unless all required inforn requests please call 1-855-4 for non-urgent requests and f	01-4262.	29.				



Epidiolex® Prior Authorization Request Form

Member Information (required)			Pr	Provider Information (required)				
Member Name:			Provider Name	Provider Name:				
Insurance ID#:			NPI#:			Specialty:		
Date of Birth:			Office Phone:					
Street Address:			Office Fax:					
City:	State:	Zip:	Office Street A	Office Street Address:				
Phone:		I	City:	City: State: Zip:			Zip:	
		Medication	Information (re	equired)				
Medication Name:			Strength:				orm:	
☐ Check if requesting	g brand		Directions for	Directions for Use:				
☐ Check if request is for continuation of therapy								
		Clinical In	formation (requ	iired)				
Select the diagnos	sis below:							
Seizures associ	ated with Drave	et syndrome						
Seizures associ	ated with Lenno	ox-Gastaut syndrome (L	,					
Other diagnosis	:		[0	CD-10 Co	de(s):			
Clinical information	n:							
Is Epidiolex prescri	bed by or in co	nsultation with a neurolo	ogist? 🗆 Yes 🗅 N	0				
Are there any other com this review?	nments, diagnoses	s, symptoms, medications t	ried or failed, and/or ar	ny other info	ormation t	the physicial	n feels is important to	
Please note: This		nied unless all required inform						

For urgent or expedited requests please call 1-855-401-4262.



Mon-Sat: 7am to 7pm Central

Evrysdi[™] Prior Authorization Request Form (Page 1 of 3) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)				
Memb	er Name:			Provider Name:			
Insura	ance ID#:			NPI#: Specialty:			
Date o	of Birth:			Office Phone:			
Street	Address:			Office Fax:			
City:		State:	Zip:	Office Street Address:			
Phone:			City:	State:	Zip:		
		М	edication Infor	mation (required)			
Medic	ation Name:			Strength: Dosage Form:			
☐ Che	eck if requesting br	and		Directions for Use:			
☐ Che	eck if request is for	continuation of thera					
			Clinical Inform	ation (required)			
	t the diagnosis be						
□ Spinal muscular atrophy (SMA): Type □ Other diagnosis: ICD-10 Code(s):							
- Oil	nei diagnosis			1CD-10 Code(s)		
1.	 Neurologist wit 		ribed by or in consultationsis and treatment of SI		ing specialis	sts:	
2.	How many SMN2 o	copies?					
	Homozygous gCompound het	gene deletion or mutatio	chromosomes 5q result in (e.g., homozygous del g., deletion of SMN1 exc	etion of exon 7 at locus		[allele 2])	
4.	ls the patient depe	ndent on invasive ventil	ation or tracheostomy? [□ Yes □ No			
5.	Is the patient depe	ndent on use of non-inv	asive ventilation beyond	use for naps and nightt	ime sleep?	⊐ Yes □ No	
	a board-certified ne ☐ Hammersmith ☐ Hammersmith ☐ Upper Limb Mo ☐ Children's Hos	eurologist? Functional Motor Scale Infant Neurological Exa odule (ULM) Test (Non	Expanded (HFMSE) m (HINE) (infant to early ambulatory) ant Test of Neuromuscul	rchildhood)		establish baseline motor ability b	
	ls the patient on co □ Yes □ No	oncomitant chronic survi	val motor neuron (SMN)	modifying therapy for th	ne treatment	of SMA (e.g., Spinraza)?	
8.	Has the patient pre	eviously received gene r	eplacement therapy for t	the treatment of SMN (e	.g., Zolgens	ma)? 🗖 Yes 🗖 No	



Evrysdi[™] Prior Authorization Request Form (Page 2 of 3) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

9.	in W hi □	patient has previously received gene therapy for the treatment of SMA (e.g., Zolgensma), provider to attests that there has been an adequate response to gene therapy (e.g., sustained decrease in at least one motor test score over a period of 6 months) or orsening in clinical status since receiving gene therapy as demonstrated by a decline of minimally clinical important difference fro ighest score achieved on one of the following exams: I HFMSE: decline of at least points on kicking and points on any other milestones (excluding voluntary grasp) I HINE-2: decline of at least points I CHOP INTEND: decline of at least points
Wh	at is at is Titr Pat Red	ty limit requests: the quantity requested per DAY? the quantity requested per DAY? the reason for exceeding the plan limitations? ration or loading dose purposes tient is on a dose-alternating schedule (e.g., one tablet in the morning and two tablets at night, one to two tablets at bedtime) quested strength/dose is not commercially available her:
		orization:
1f ti	Pro	s a reauthorization request, answer the following: ovide documentation of positive clinical response to therapy (e.g., chart notes, laboratory values) from pretreatment baseline status demonstrated by the most recent results (less than 1 month prior to reauthorization request) from one of the following exams: One of the following HINE-2 milestones Improvement or maintenance of previous improvement of at least a 2-point (or maximal score) increase in ability to kick Improvement or maintenance of previous improvement of at least a 1-point increase in any other HINE-2 milestone (e.g.,
		 head control, rolling, sitting, crawling, etc.), excluding voluntary grasp Patient exhibited improvement, or maintenance of previous improvement in more HINE motor milestones than worsening, from pretreatment baseline (net positive improvement) Patient has achieved and maintained any new motor milestones when they would otherwise be unexpected to do so (e.g., sit unassisted, stand, walk)
		One of the following HFMSE milestones Improvement or maintenance of a previous improvement of at least a 3-point increase in score from pretreatment baseline Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so (e.g., sit unassisted, stand, walk)
		One of the following ULM test milestones Improvement or maintenance of a previous improvement of at least a 2-point increase in score from pretreatment baseline Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so (e.g., sit unassisted, stand, walk)
		One of the following CHOP-INTEND milestones Improvement or maintenance of a previous improvement of at least a 4-point increase in score from pretreatment baseline Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so (e.g., sit unassisted, stand, walk)
		One of the following MFM-32 milestones Improvement or maintenance of a previous improvement of at least a 3-point increase in score from pretreatment baseline Patient has achieved and maintained any new motor milestone from pretreatment baseline when they would otherwise be unexpected to do so (e.g., sit unassisted, stand, walk)
2.	ls t	he patient dependent on invasive ventilation or tracheostomy? Yes No
3.	ls t	he patient dependent on the use of non-invasive ventilation beyond use for naps and nighttime sleep? Yes No



Evrysdi[™] Prior Authorization Request Form (Page 3 of 3) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

4.	Is the requested medication prescribed by or in consultation with a neurologist with expertise in the diagnosis and treatment of SMA? — Yes — No
5.	Is the patient is receiving concomitant chronic survival motor neuron (SMN) modifying therapy for the treatment of SMA (e.g., Spinraza)? Yes No
6.	Has the patient previously received gene replacement therapy for the treatment of SMA (e.g., Zolgensma)? □ Yes □ No
7.	Was there inadequate response to gene therapy (e.g., sustained decrease in at least one motor test score over a period of 6 months)? If so, submit medical records (e.g., chart notes) documenting the inadequate response to gene therapy.
	here any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to review?
Pleas	se note: This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262

For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.



Genitourinary smooth muscle relaxants Prior Authorization Request Form

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:	Provider Name:		
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:			City:	State:	Zip:	
		Medication	nformation (red	quired)		
Medication Name:			Strength:	1	Dosage Form:	
☐ Check if requesting	g brand		Directions for Us	se:		
☐ Check if request is		on of therapy				
		Clinical Inf	ormation (requi	red)		
What is the patien	t's diagnosis	for the medication beir				
panen	ar a anagmasia		.g requeeteur (ar	,		
ICD-10 Code(s) [N	//andatory]: _					
Medication history	y:					
Has the patient had	d a 30-day trial	of oxybutynin or oxybuty	nin extended-releas	e (ER)? U Ye s	s □ No	
-	-	ests, also answer the fo	_			
Does the patient ha	ave a diagnosi	s which confirms a difficu	Ity in swallowing?	I Yes □ No		
Quantity limit requestion What is the quantity		ar MONTH?				
·		ng the plan limitations?				
□ Titration or load	ing dose purpo	oses				
		g schedule (e.g., one tab	let in the morning an	nd two tablets a	t night, one to two	
tablets at bedtin	,	ot commercially available				
☐ Other:	.g, 4000 10 11	or commercially available				
Are there any other com this review?	nments, diagnos	es, symptoms, medications to	ried or failed, and/or any	other information	n the physician feels is important to	
Please note: This	request may be d	enied unless all required inform	ation is received.			

For urgent or expedited requests please call 1-855-401-4262.



GLP-1 Agonists Prior Authorization Request Form

Member Information (required)			Provider Information (required)					
Member Name:			Provider Nam	Provider Name:				
Insurance ID#:			NPI#:		Specialty:			
Date of Birth:			Office Phone	:				
Street Address:			Office Fax:					
City:	State:	Zip:	Office Street					
Phone:		L	City:	State:	Zip:			
		Medication	Information	(required)				
Medication Name:			Strength:					
☐ Check if requesti	ng brand		Directions for	· Use:				
☐ Check if request	is for continuatio	on of therapy						
		Clinical Ir	nformation (re	equired)				
Select the diag	nosis below:							
☐ Type 2 diabe								
Other diagno	osis:		ICD-10 Co	ICD-10 Code(s):				
Quantity limit r	•	I per MONTH?						
·	• •	eding the plan limitati						
☐ Titration or lo	oading dose pu	ırposes						
		ating schedule (e.g., or	ne tablet in the m	orning and two ta	ablets at night, one to two			
tablets at bed	,	s not commercially ava	ilahla					
		s not commercially ava						
Are there any other of this review?	comments, diagno	ses, symptoms, medications	tried or failed, and/or	any other informatio	on the physician feels is important to			
Please note: T	his request may be	denied unless all required infor						

For urgent or expedited requests please call 1-855-401-4262.



Gralise® & Horizant® Prior Authorization Request Form

Member Information (required)				Provider Information (required)				
Member Name:			Provider Name	Provider Name:				
Insurance ID#:			NPI#:		Specialty:			
Date of Birth:	Date of Birth:			Office Phone:				
Street Address:			Office Fax:					
City:	City: State: Zip:			Address:				
Phone:			City:	State:	Zip:			
		Medicatio	n Information (required)				
Medication Nam	ne:		Strength:		Dosage Form:			
☐ Check if requ	esting brand		Directions for	Use:				
☐ Check if requ	est is for continuatio r	n of therapy						
		Clinical	Information (red	quired)				
Select the di	iagnosis below:							
□ Moderate	to severe primary	restless leg syndror	me (RLS) [Horizant	only]				
□ Neuropatl	hic pain associated	d with postherpetic n	euralgia (PHN)					
Other diag	gnosis:		ICD-10 Co	_ ICD-10 Code(s):				
	severe primary F							
		failure (to a minimun days? 🗖 Yes 🗖 No	n of a 90 day trial),	contraindicatio	n, or intolerance to ropinirole			
Neuropathic	pain associated	with PHN:						
		failure (to a minimun in the past 180 days		contraindicatio	n, or intolerance to an			
Are there any oth this review?	her comments, diagnos	es, symptoms, medicatior	ns tried or failed, and/or a	any other information	on the physician feels is important to			
Please note:		enied unless all required inf						



Growth Hormones Prior Authorization Request Form (Page 1 of 3)

DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)				
Member Name:		· · ·	Provider	Provider Name:			
Insurance ID#:			NPI#:	NPI#: Specialty:			
Date of Birth:			Office Ph	ione:			
Street Address:			Office Fa	x:			
City:	State:	Zip:	Office Str	Office Street Address:			
Phone:			City:		State:		Zip:
		Medication	Informatio	n (required)			
Medication Name:				Strength: Dosage Form:			orm:
☐ Check if requesting	g brand		Direction	s for Use:			
☐ Check if request is	for continuation	of therapy					
		Clinical In	formation	(required)			
☐ Small for gestation	deficiency in childrent to chronic renal in the to panhypopituitate to Prader-Willi sylature in childrent en	en nsufficiency rism	,				
☐ Turner's syndrome For Adults (18 years ☐ Growth hormone of	of age or older):						
□ Panhypopituitarisr	n						
☐ Prader-Willi syndrome ☐ Other diagnosis: ICD-10 Code(s):							
Contraindications/E				100-10 000			
Does the patient have trauma, or acute resp Does the patient have	e acute critical illne iratory failure?		.				Itiple accidental



Growth Hormones Prior Authorization Request Form (Page 2 of 3) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

s the requested medication prescribed by or in consultation with a pediatric endocrinologist?
Has the patient been screened for intracranial malignancy or tumor?
For growth hormone deficiency in children, also answer the following: Has the patient had an inadequate response to two (2) pharmacological growth hormone stimulation tests* with peak level below 10 ng/mL? Yes No Has the patient had an inadequate response to at least one (1) pharmacological growth hormone stimulation tests* with peak level below 10 ng/mL for a patient with defined CNS pathology, multiple pituitary hormone deficiencies, history of irradiation, or proven genetic cause? Yes No *Please note: acceptable tests include: arginine, clonidine, glucagon, insulin, and levodopa s the patient's height more than 3 standard deviations (SDs) below the mean for same age and gender? Yes No s the patient's height more than 2 SDs below the mean for same age and gender AND the patient has decreased growth velocity more than 1 SD below the mean for the same age and gender? Yes No s the patient's growth velocity measured 2 SDs below the mean over one year or 1.5 SDs below the mean sustained over 2 years for the same age and gender? Yes No Has the patient had an inadequate response to two (2) pharmacological growth hormone stimulation tests* with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient with peak level below 10 ng/mL for a patient wit
Has growth hormone deficiency been confirmed with provocative test and/or IGF-1 levels?
Has the patient had an inadequate response to two (2) pharmacological growth hormone stimulation tests* with peak level below 10 ng/mL?
Has the patient had an inadequate response to at least one (1) pharmacological growth hormone stimulation test* with peak level below 10 ng/mL for a patient with defined CNS pathology, multiple pituitary hormone deficiencies, history of irradiation, or proven genetic cause?
below 10 ng/mL for a patient with defined CNS pathology, multiple pituitary hormone deficiencies, history of irradiation, or proven genetic cause?
s the patient's height more than 3 standard deviations (SDs) below the mean for same age and gender? \(\begin{align*} \text{Yes} \text{No} \\ s the patient's height more than 2 SDs below the mean for same age and gender AND the patient has decreased growth velocity more han 1 SD below the mean for the same age and gender? \(\begin{align*} \text{Yes} \text{No} \\ s the patient's growth velocity measured 2 SDs below the mean over one year or 1.5 SDs below the mean sustained over 2 years for he same age and gender? \(\begin{align*} \text{Yes} \text{No} \\ \text{Have other causes of growth failure been ruled out (e.g., hypothyroidism, chronic systemic disease, skeletal disorders, malnutrition)? \(\begin{align*} \text{Yes} \text{No} \\ \text{No} \end{align*}
s the patient's height more than 2 SDs below the mean for same age and gender AND the patient has decreased growth velocity more han 1 SD below the mean for the same age and gender?
han 1 SD below the mean for the same age and gender? \(\begin{align*} \text{Yes} \end{align*} \text{No} \\ s the patient's growth velocity measured 2 SDs below the mean over one year or 1.5 SDs below the mean sustained over 2 years for he same age and gender? \(\begin{align*} \text{Yes} \end{align*} \text{No} \\ Have other causes of growth failure been ruled out (e.g., hypothyroidism, chronic systemic disease, skeletal disorders, malnutrition)? \(\begin{align*} \text{Yes} \end{align*} \text{No} \end{align*} \)
he same age and gender?
malnutrition)?
For growth failure due to chronic ronal incufficiency, also answer the following:
for growth failure due to chromic renarmisanticiency, also answer the following.
Has the patient's nutritional status been optimized and metabolic abnormalities been corrected? Yes No
Has the patient had a kidney transplant? ☐ Yes ☐ No
s the patient's height less than the 3 rd percentile?
s the patient's growth velocity measured over 1 year > 2 standard deviations below the mean for same age and gender? \(\Quad Yes \) No
For growth failure due to panhypopituitarism or Prader-Willi syndrome, also answer the following: Has the patient's diagnosis of panhypopituitarism or Prader-Willi syndrome been confirmed by appropriate genetic esting? No
s the diagnosis of panhypopituitarism caused by cranipharyngioma surgery? Yes No
Does the patient have severe obesity, history of upper airway obstruction or sleep apnea, or severe respiratory mpairment? Type No
s the patient's height more than 2 standard deviations below the mean for same age and gender? Yes No
For idiopathic short stature, also answer the following:
s the patient's height more than 2.25 standard deviations below the mean? Yes No
s the patient's predicted height less than or equal to 65 inches for male or less than or equal to 60 inches for females? 🗖 Yes 🗖 No
For short stature homeobox-containing gene (SHOX) deficiency or Noonan syndrome, also answer the following: s the patient's height more than 3 standard deviations (SDs) below the mean for same age and gender? Yes No s the patient's height more than 2 SDs below the mean for same age and gender AND the patient has decreased growth velocity more han 1 SD below the mean for the same age and gender? Yes No
s the patient's growth velocity measured 2 SDs below the mean over one year or 1.5 SDs below the mean sustained over 2 years for he same age and gender?
For small for gestational age (SGA), also answer the following:
Did the patient have post-natal growth failure at one year?
s the patient below the 5 th percentile for height?
Was the patient's birth weight or length at least 2 standard deviations below the mean for gestational age? Yes No
For Turner's syndrome, also answer the following:
Has the patient's diagnosis of Turner's syndrome been confirmed by chromosome analysis?



Growth Hormones Prior Authorization Request Form (Page 3 of 3) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

For Adult Pa	atients (18 years of age or older):						
Is the reques	Is the requested medication prescribed by or in consultation with an endocrinologist? Yes No						
For growth	hormone deficiency in adults, also answer the following:						
Has growth h	normone deficiency been confirmed with two provocative tests and IGF-1 levels? Yes No						
Has the patie	ent been screened for intracranial malignancy or tumor?						
Are there any o	ther comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to						
Please note:	This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.						

61



Serostim® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:	<u>l</u>	I	City:	State:	Zip:	
		Medication Info	ormation (required	1)		
Medication Name:			Strength:	<u></u>	Dosage Form:	
☐ Check if requesting	brand		Directions for Use:			
☐ Check if request is	for continuation of					
		Clinical Infor	mation (required)			
Select the diagnosis below: HIV infection/AIDS wasting Other diagnosis: ICD-10 Code(s): Clinical information: Is Serostim prescribed by or in consultation with an infectious disease specialist? Yes No Has the patient tried and had an inadequate response or intolerance to dronabinol or megestrol? Yes No Is the patient currently receiving treatment with antiretrovirals? Yes No Does the patient have acute critical illness due to complications following open heart surgery, abdominal surgery, multiple accidental trauma, or those with acute respiratory failure? Yes No Has the patient been screened to verify the absence of any active malignancy? Yes No Does the patient have active proliferative or severe non-proliferative diabetic retinopathy? Yes No Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?						
Please note: This	request may be denied	I unless all required information	on is received.			

For urgent or expedited requests please call 1-855-401-4262.



Zorbtive® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Ac	ldress:		
Phone:	I	1	City:	State:	Zip:	
Medication Information (required)						
Medication Name:			Strength:		Dosage Form:	
☐ Check if requesting	brand		Directions for U	se:		
☐ Check if request is t	for continuation of th	erapy				
		Clinical Inform	mation (requir	red)		
Select the diagnosis below: Short bowel syndrome Other diagnosis:						
Is the patient receiving Does the patient has accidental trauma, of	ed by or in consultation of specialized nutrition of acute critical illness or acute respiratory for	on with a gastroentero ional support (i.e., paress due to complications ailure?	enteral nutrition) s following open	?	bdominal surgery, multiple	
Are there any other conthis review?	nments, diagnoses, syn	nptoms, medications tried	or failed, and/or ar	ny other information	n the physician feels is important to	
Please note: This	request may be denied u	nless all required information	n is received			

For urgent or expedited requests please call 1-855-401-4262.



Lindane shampoo, Ovide® (malathion), NatrobaTM (spinosad), Sklice® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)				
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street	Address:			
Phone:		l	City:	State:		Zip:	
		Medication Inf	ormation	(required)			
Medication Name:			Strength:				
☐ Check if request			Directions for Use:				
□ Check if request	is for continuati	on of therapy					
		Clinical Info	rmation (re	equired)			
Medication his	story:						
		d failure, contraindication, 0 days? ☐ Yes ☐ No	or intolerance	e to a permethrir	or pyrethr	ins-piperonyl	
butoxide produc	or in the past s	o days! La res La No					
Are there any other cothis review?	omments, diagnos	ses, symptoms, medications tried	or failed, and/or	any other information	on the physicia	an feels is important to	
		denied unless all required information de requests please call 1-855-401-4.					



Hemangeol[™] Prior Authorization Request Form

Member Information (required)			Provider Information (required)				
Member Name:		Provider Name	Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth: Street Address:			Office Phone:				
			Office Fax:				
City: State: Zip:			Office Street A	Address:			
Phone:			City:	State:	Zip:		
		Medication	Information	(required)			
Medication Nar	me:		Strength:		Dosage Form:		
☐ Check if requ	uesting brand		Directions for	Use:			
☐ Check if requ	uest is for continuation	of therapy					
		Clinical Ir	nformation (red	quired)			
Select the d	liagnosis below:						
Proliferat	ing infantile heman	gioma requiring syste	mic therapy				
Other dia	agnosis:		ICD-10 Co	ode(s):			
Clinical info	ormation:						
		ms (kg) or greater?					
•		or a history of broncho	•				
•	•	dia (less than 80 bea	•				
•	•	han first-degree heart	•		ure? 🛘 Yes 🗎 No		
-	•	essure less than 50/30	•	i □ No			
Does the pa	tient have pheochro	omocytoma? Yes	□ No				
Are there any ot this review?	ther comments, diagnose	es, symptoms, medications	tried or failed, and/or	any other informatio	on the physician feels is important to		
Please note:		enied unless all required infor					



Hepatitis C Prior Authorization Request Form (Page 1 of 2)

		JRE USE. FORMS ARE UP				
Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#: Specialty:			
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:			City:	State:		Zip:
		Medication Info	rmation (required)			
Medication Name:			Strength: Dosage Form:			orm:
☐ Check if requesting	brand		Directions for Use:			
☐ Check if request is	for continuation of the	rapy				
		Clinical Inforr	nation (required)			
Select the diagnosis	below:					
☐ Hepatitis C virus ir						
			ICD-10 Cod	le(s):		
Clinical information:						
Document the patient	's genotype: 's weight:	 Ka				
		ə lers may be asked to pro	vide documenatation):			
Liver biopsy con	firming a Metavir score	of F2 or greater. List F2	score			
Serum aspartate	e aminotransferase (AS	Γ)-to-platelet ratio index ((APRI) score of 1.5 or gr	eater? List	APRI score	
☐ Fibroscan score	of 7.1 kPa or greater. L	ist fibroscan score	- O :- f f - - -		.4	Baardana Iddaa
		manifestations of hepatiti			itoimmune a	isorders, kidney
disease, severe						
Patient is curren	ıtly pregnant. How was ı	oregnancy confirmed?				
Does the patient have	e cirrhosis? Yes	No				
Does the patient have	compensated liver dise	ease (Child-Pugh A)? 📮				
	e decompensated liver d nt naïve?	isease (Child-Pugh B or	C)? Yes No			
· ·		ואס ug and alcohol free for th	e past 6 months? DYe	s 🗆 No		
		n, does the patient have			days prior to	o initiation of
therapy and will receive	ve a monthly pregnancy	test during treatment?	☐ Yes ☐ No			
_	~	elpatasvir, also answe				
		e to ribavirin?				
Has the patient had prioir treatment failure with sofosbuvir or NS5A-based tre			ased treatment? Yes	□ No		
Is the patient taking P glycoprotein (P-gp) inducers? Yes No						
Is the patient taking anticancers (e.g., topotecan)? Yes No Is the patient taking moderate to potent CYP inducers (e.g., rifampin, St. John's wort, carbamazepine, pho						
oxcarbazepine)?	•	inducers (e.g., mampin,	St. John's Wort, Carbama	azepine, pri	enytoin, prie	nobarbitai,
. ,		ofosbuvir, also answer	the following:			
	taking any of the follow		5			
Anticonvulsants	(e.g., carbamazepine, d	oxcarbazepine, phenobai	rbital, phenytoin)			ing HIV regimens
	P-gp) inducers (e.g., rifa			☐ Antica	aners (e.g., t	opotecan)
HIV antiretrovira	ıls (e.g., tipranavir/ritona	IVIF)				



Hepatitis C Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

For Mavyret, also answer the following:
Select if the patient has been previously treated with a regimen containing the following (select all that applies):
□ An HCV NS5A inhibitor
□ An NS3/4A protease inhibitor (PI) □ Interferon (including pegylated formulations), ribavirin, and/or Sovaldi (sofosbuvir)
For Sovaldi, also answer the following:
Select if the patient will use Sovaldi in combination with the following:
Pegylated interferon and ribavirin
☐ Ribavirin
Does the patient have severe renal impairment (eGFR < mL/min/1.73 m²)? ☐ Yes ☐ No
Does the patient have end-stage renal disease? ☐ Yes ☐ No
Does the patient have hepatocellular carcinoma that meets criteria for liver transplant? Yes No
For Vosevi, also answer the following:
Has the patient been previously treated with a regimen containing an NS5A inhibitor? ☐ Yes ☐ No
Has the patient been previously treated with a regimen containing Sovaldi (sofosbuvir) without an NS5A inhibitor? Yes No
For Zepatier, also answer the following:
Has the patient been tested for the presence of NS5A resistance-associated polymorphisms? ☐ Yes ☐ No
If yes to the above question, does the patient have baseline NS5A polymorphisms? Yes No
Does the patient have moderate to severe hepatic impairment?
Has the patient failed the 2-drug regimen of peginterferon alfa and ribavirin? ☐ Yes ☐ No
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to his review?
IIIS TEVIEW!
_
Please note: This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.



Brand Name narcotics Prior Authorization Request Form (Page 1 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)				Provider Information (required)			
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Addre	ess:			
Phone:	<u> </u>		City:	State:		Zip:	
		Medication I	nformation (requ	ired)			
Medication Name:			Strength:	,	Dosage Fo	orm:	
☐ Check if requesting	g brand		Directions for Use:				
☐ Check if request is	for continuation of the	nerapy					
		Clinical Inf	ormation (require	d)			
Medication history:	trial and fallers (-1)		and the second		Non DN		
Clinical information:	•	ast a 30 day trial) of a g	eneric narcotic in the pa	ist 90 days? L	I Yes U No		
		r in the past 365 days?	□ Yes □ No				
•	·	inal illness? 🛘 Yes 🚨					
•	-		.g., sickle cell anemia, e	etc)? 🗖 Yes 🗆	⊒ No		
If yes, please list the	-						
		with significant pain?	l Yes □ No				
		to the lowest effective					
If yes , please provide	documentation:						
Reauthorization:							
If this is a reauthoriz	=	-	10 D.V. D.V.				
Is the prescriber main If yes , please provide		ervative, effective treatr	nent? Li Yes Li No				
ii yes , picase provide	documentation.						
Quantity limit reques							
What is the patient's	s diagnosis for the m	edication being reque		Code(s)			
What is the quantity re	equested per MONTH	?					
What is the reason f	or exceeding the pla						
☐ Titration or loading☐ Patient is on a dos	g dose purposes se-alternating schedule	e (e.g., one tablet in the	morning and two table	ts at night one	to two tablets	s at bedtime)	
Requested strengt	th/dose is not commer	cially available		at mymt, one	the tablete	o at bouning)	
☐ Other:							



Brand Name narcotics Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.



Hydrocodone-acetaminophen (APAP) Products Prior Authorization Request Form (Page 1 of 2)

	DO NOT COPY FOR	FUTURE USE. FORMS ARE	UPDATED FREQU	JENTLY AND MAY B	E BARCODED	
Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street A	Address:		
Phone:			City:	State:	Zip:	
		Medication In	formation	(*************************************		
Medication Name:		Medication in	Strength:	(required)	Dosage Form:	
☐ Check if requesting	g brand		Directions for	Use:		
☐ Check if request is	•	therapy				
		Clinical Info	rmation (red	quired)		
Medication histor	v·			quireu)		
Has the patient had a history of a 60 day trial (in the past 90 days) with one of the following generics listed below? • Hydrocodone-APAP 5-325 • Hydrocodone-APAP 10-325 Clinical information:						
•	-	cancer in the past 365 d	•	l No		
-	=	a terminal illness? Ye		!!	D.V. D.N.	
If yes , please list the	· <u></u>	ciated with significant pa	iin (e.g., sickie (ceii anemia, etc)?	LI Yes LI NO	
	ave an <u>injury</u> assoc	iated with significant pai	n? 🗆 Yes 🗅 N	No		
Have efforts been made to taper the patient to the lowest effective dose?						
Is the prescriber ma	aintaining the most	answer the following: conservative, effective				



Hydrocodone-acetaminophen (APAP) Products Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Are there any of this review?	ther comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to
Please note:	This request may be denied unless all required information is received.



Morphine Equivalent Dose (MED) Limit Prior Authorization Request Form

Member Information (required)			Pr	Provider Information (required)			
Member Name:			Provider Name:	Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Ad	Office Street Address:			
Phone:			City:	State:	Zip:		
		Medication I	nformation (re	equired)			
Medication Name:			Strength:	equil out	Dosage Form:		
☐ Check if requesting	brand		Directions for U	se:			
☐ Check if request is		herapy					
		Clinical Inf	ormation (requ	ired)			
Clinical information	on:		<u> </u>	<u> </u>			
Does the patient ha	ve a diagnosis of ca	ancer in the past 365	days? 🗆 Yes 🗅 I	No			
Does the patient ha	ve a diagnosis of a	terminal illness?	Yes □ No				
Does the patient ha	ive an <u>illness</u> assoc	iated with significant	pain (e.g., sickle ce	ell anemia, etc)?	☐ Yes ☐ No		
If yes , please list th	e diagnosis:						
•		ated with significant p		•			
• •	-						
		atient to the lowest ef					
ii yes , piease provi	de documentation: <u>.</u>						
Reauthorization:							
		nswer the following					
-	~	conservative, effectiv					
if yes , piease provid	de documentation: _.						
Are there any other co	mments, diagnoses, sy	mptoms, medications tr	ied or failed, and/or an	y other informatio	n the physician feels is important to		

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262.



Opioid Naïve Prior Authorization Request Form

Meml		ation (required)	Provider Information (required)				
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Ad	dress:			
Phone:			City:	State:	Zip:		
		Medication	Information (re	equired)			
Medication Name:			Strength:	~(Dosage Form:		
☐ Check if requesting	g brand		Directions for Us	se:			
☐ Check if request is	for continuatio r	of therapy					
		Clinical In	formation (requi	ired)			
Clinical information	on:						
Does the patient ha	ave a diagnosis	of cancer in the past 369	5 days? ☐ Yes ☐ N	No			
Does the patient ha	ave a diagnosis	of a terminal illness?	Yes □ No				
Does the patient ha	ave an <u>illness</u> a	ssociated with significant	t pain (e.g., sickle ce	II anemia, majo	r surgery, etc)?		
If yes , please list th	ne diagnosis:	·					
Does the patient ha	ave an <u>injury</u> as	sociated with significant	pain? 🗆 Yes 🗅 No	•			
If yes , please list th	ne diagnosis:						
Have efforts been r	made to taper th	ne patient to the lowest e	effective dose? 🗖 Ye	es 🗆 No			
If yes , please provi	de documentat	ion:					
Are there any other co this review?	mments, diagnos	es, symptoms, medications t	tried or failed, and/or an	y other informatio	on the physician feels is important to		
Please note: This	e request may be d	enied unless all required inform	mation is received				



Long Acting and Short Acting Opioid Prior Authorization Request Form

Ме	Member Information (required)			Provider Information (required)			
Member Name:			Provider Name	Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:	I			
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Ac	ddress:			
Phone:			City:	State:	Zip:		
		Medication	Information (re	equired)			
Medication Name	e:	modification.	Strength:		Dosage Form:		
☐ Check if reque	esting brand		Directions for U	Jse:			
	est is for continuation	of therapy					
		Clinical Ir	nformation (requ	uired)			
Clinical inform	nation:		` '	,			
		of cancer in the past 3	365 davs? □ Yes □	l No			
	=	of a terminal illness?	-				
•	ŭ	ssociated with significa		cell anemia etc)?	□ Yes □ No		
•	st the diagnosis:						
Does the patier	nt have an <u>injury</u> ass	sociated with significar	nt pain? 🛚 Yes 🗖 N	lo			
-	st the diagnosis:		·				
Have efforts be	en made to taper th	ne patient to the lowes	t effective dose? 🗖	Yes □ No			
lf yes , please p	provide documentati	on:					
Reauthorization							
	-	st, answer the follow	_				
•	-	ost conservative, effec	ctive treatment?	′es □ No			
If yes , please p	provide documentati	on:					
re there any other iis review?	comments, diagnoses	s, symptoms, medications	tried or failed, and/or an	ny other information the	ne physician feels is important		

This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262. Please note:



Esbriet® & Ofev® Prior Authorization Request Form

Member Information (required)			Pro	Provider Information (required)			
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street A	ddress:			
Phone:	I	I	City:	State:		Zip:	
		Medication	Information (re	quired)			
Medication Name:		-	Strength:		Dosage F	orm:	
☐ Check if reques	ting brand		Directions for U	Jse:			
□ Check if reques	t is for continuatior	of therapy					
		Clinical Inf	formation (requi	ired)			
Select the diagr	nosis below:						
☐ Idiopathic pul	monary fibrosis (I	PF)					
□ Other diagnor	sis:		ICD-1	0 Code(s):			
Clinical informa	ation:						
Does the patient days? Yes		al capacity (FVC) greate	er than or equal to 5	0% of predicted i	n the last 6	0	
•		ribed by or in consultation	on with a pulmonolo	gist? 🗆 Yes 🚨	No		
<u> </u>	·	·	· · · · · · · · · · · · · · · · · · ·	<u> </u>			
Are there any other on this review?	comments, diagnoses	s, symptoms, medications t	tried or failed, and/or ar	ny other information	the physicia	n feels is important t	



Actemra® Prior Authorization Request Form (Page 1 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:	<u>l</u>	-L	City:	State:	Zip:	
		Medication Info	ormation (required)			
Medication Name:			Strength:		Dosage Form:	
☐ Check if requesting	brand		Directions for Use:			
	for continuation of the	erapy				
		Clinical Inform	nation (required)			
Clinical Information (required) Select the diagnosis below: Active polyarticular juvenile idiopathic arthritis (pJIA) Active systemic juvenile idiopathic arthritis (sJIA) Chimeric antigen receptor (CAR) T cell-induced severe or life-threatening cytokine release syndrome (CRS) Moderately to severely active rheumatoid arthritis (RA) Temporal arteritis or giant cell arteritis (GCA) Systemic sclerosis-associated interstitial lung disease Other diagnosis: ICD-10 Code(s): Clinical information: Select if Actemra is prescribed by or in consultation with one of the following specialists: Allergist/immunologist Rheumatologist Other Will Actemra be used in combination with another biologic agent or targeted immunomodulator? Yes No						
Has the patient had a	•	arthritis (pJIA), also and to, intolerance to, or con		ore non-bio	logic disease modifying anti-	
For active systemic	juvenile idiopathic art	hritis (sJIA), also answ		tlie non-s	teroidal anti-inflammatory drugs	
(NSAIDs), corticostero		of intolerance to at least	one oral systemic agen	t [i.e., fiori-s	teroidal anti-lillianimatory drugs	
_		toid arthritis (RA), also				
· ·	n inadequate response ARDs)? ☐ Yes ☐ No	to, intolerance to, or con	traindication to one or m	ore non-bio	logic disease modifying anti-	
For temporal arteritis Has the patient had a						



Actemra® Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?								
Please note:	This request may be denied unless all required information is received.							



Adbry® Prior Authorization Request Form

Member Information (required)				Provider Information (required)			
Member Name:			Provider Name	e:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street A	Address:			
Phone:	<u> </u>	l	City:	State:	Zip:		
		Medication	Information (re	equired)	<u> </u>		
Medication Name:			Strength:		Dosage Form:		
☐ Check if requesting			Directions for	Use:			
☐ Check if request is	for continuatior						
		Clinical In	nformation (requ	ıired)			
Select the diagnosis							
Atopic dermatitis (cOther diagnosis:	describe severity	level)	ICD	0-10 Code(s):			
Clinical information:				.,			
Select if the requested Dermatologist	d medication is p Allerg	rescribed by or in consulti ist/Immunologist	tation with one of the fo	ollowing specialists	:		
Medication history:	<u>.</u>						
Has the patient have a	a documented 14	l in combination with anot 4-day trial of a topical cor	ticosteroid, pimecrolim	us cream, tacrolim			
·							
Are there any other com this review?	ments, diagnoses	s, symptoms, medications	tried or failed, and/or ar	ny other information	the physician feels is important to		
							
Please note: This r	aguast may ba dar	nied unless all required infor	mation is received				

For urgent or expedited requests please call 1-855-401-4262.



Cibinqo™ Prior Authorization Request Form

DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Memb	er Informat	ion (required)	Pro	ovider Info	rmation (required)
Member Name:			Provider Name	e:	
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone:		
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street A	ddress:	
Phone:			City:	State:	Zip:
		Medication	Information (re	equired)	<u> </u>
Medication Name:			Strength:		Dosage Form:
☐ Check if requesting			Directions for U	Jse:	
☐ Check if request is	for continuation o				
		Clinical In	formation (requ	ired)	
Select the diagnosis		wal)			
		evel)	ICD	-10 Code(s):	
Clinical information: Select if the requested Dermatologist		scribed by or in consulta	ation with one of the fo		:
Has the patient have a	documented 14-c	combination with anoth	icosteroid, pimecrolimu	-	odulator?
Are there any other comithis review?	ments, diagnoses, s	symptoms, medications t	tried or failed, and/or an	y other information	the physician feels is important to

<u>Please note</u>: This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Cimzia® Prior Authorization Request Form (Page 1 of 2)

Mem	Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:	:			
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street	Address:			
Phone:			City:	State:		Zip:	
		Madiaation				·	
Medication Name:		Medication	Information (required)	Decemb	o rm.	
			Strength:		Dosage F	om.	
☐ Check if requesti			Directions for	Use:			
☐ Check if request	is for continuatio		f 1:				
		Clinical in	formation (req	uired)			
Select the diagnos							
☐ Active ankylosing							
☐ Active psoriatic a							
☐ Moderate to seve		•					
☐ Moderately to se	•						
☐ Moderately to se	•						
Active non-radioOther diagnosis:		idyloartifitis	ICI	D-10 Code(s):			
			ICL	J-10 Code(s)			
Clinical informatio		properited by ar in consults	ation with one of the f	allowing anasialists			
□ Dermatologist		prescribed by or in consulta enterologist	auon with one or the r eumatologist	Ollowing specialistsOther	5.		
•		d in combination with anoth	•		odulator?	Yes □ No	
-		also answer the following		argotou mmanome			
		sponse to, intolerance to, o		one or more non-st	teroidal anti-in	nflammatory drugs	
(NSAIDs)? • Yes	□ No	·					
		enswer the following:					
		sponse to, intolerance to, o		methotrexate? 🔲 `	Yes 🛚 No		
	-	aque psoriasis, also answ					
		sponse to, intolerance to, o				st one of the	
sulfasalazine)?		ore oral systemic treatments	s (i.e., methotrexate, o	cyclosporine, acitre	etin,		
For moderately to	severely active (Crohn's disease, also ans	wer the following:				
		esponse to, intolerance to, o trexate)? ☐ Yes ☐ No	or contraindication to	one or more immur	nosuppressive	e agents (e.g.,	
For moderately to	severely active r	heumatoid arthritis, also	answer the following	ıg:			
Has the patient had rheumatic drugs (DI		esponse to, intolerance to, o	or contraindication to	one or more non-bi	iologic diseas	e modifying anti-	
	• .	spondyloarthritis, also an	_				
		esponse to, intolerance to, o trexate)? ☐ Yes ☐ No	or contraindication to	one or more immur	nosuppressive	e agents (e.g.,	



Cimzia® Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

<u>Please note</u>: This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Cosentyx® Prior Authorization Request Form

Men		or future use. Forms A			rmation (required)	
Member Name:			Provider Name	Provider Name:		
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
	Otata	7:		d du a a a a		
City:	State:	Zip:	Office Street A	aaress:		
Phone:			City:	State:	Zip:	
		Medication	Information (re	quired)		
Medication Name:			Strength:		Dosage Form:	
☐ Check if requesti	ng brand		Directions for U	Jse:		
☐ Check if request	<u> </u>	on of therapy				
		Clinical In	nformation (requi	ired)		
Select the diagnos	sis helow:		(loqui	ou		
☐ Active ankylosin						
☐ Active psoriatic						
☐ Moderate to sev		nsis				
☐ Active Non-radio						
☐ Active enthesitis	• .	mayroarannao				
☐ Other diagnosis:			ICD-	-10 Code(s):		
Clinical information				.,		
Select if the reques Dermatologist	ted medication is	prescribed by or in consult umatologist	her			
Will the requested r	nedication be use	ed in combination with anot	her biologic agent or ta	rgeted immunomo	dulator?	
For active ankylos	sing spondylitis,	also answer the followin	g:			
Has the patient had (NSAIDs)? ☐ Yes		esponse, contraindication, o	or intolerance to one or	more non-steroid	lal anti-inflammatory drugs	
-	•	answer the following:			- N	
· .	•	esponse, contraindication,		trexate? L Yes (⊔ No	
		oriasis, also answer the f	-			
		esponse, contraindication, on the stemic treatments (i.e., met			h at least one of the following: lazine)? ☐ Yes ☐ No	
	• .	spondyloarthritis or enth			•	
Has the patient had (NSAIDs)? \(\begin{array}{c}\Ds\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	•	esponse, contraindication, o	or intolerance to one or	more non-steroid	lal anti-inflammatory drugs	
Are there any other co	omments, diagnos	es, symptoms, medications	tried or failed, and/or an	y other information	the physician feels is important to	

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262.



Dupixent® Prior Authorization Request Form (Page 1 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address	:		
Phone:		1	City:	State:		Zip:
		Medication Inf	ormation (required	d)		
Medication Name:			Strength:	,	Dosage Fo	orm:
☐ Check if requesting	g brand		Directions for Use:			
	for continuation of th	erapy				
		Clinical Infor	mation (required)			
Select the diagnos	sis below:					
☐ Atopic dermatitis	5					
☐ Chronic rhinosin	usitis with nasal poly	posis (CRSwNP)				
■ Moderate to sev	ere asthma					
Eosinophilic eso	phagitis					
☐ Prurigo nodulari:	s					
Other diagnosis:	· ·			0-10 Code(s):	
Atopic dermatitis:						
			oid, pimecrolimus cre	eam, tacroli	imus ointm	ent, Eurisa
, ,		days? 🗖 Yes 🗖 No				
	•		logist or allergist/imm	iunologist?	⊔ Yes ∟	I NO
	sitis with nasal poly	• • •	000 ND0 D.V			
· ·	-	•	CRSwNP? Yes		00 10 1	D.V D.N.
			osteroid (INCS) withir		-	
(i.e., ENT)? \(\begin{array}{c}\) Yes		itation with an allergis	st/immunologist, pulm	onologist, (orololaryn	gologist
Moderate to sever	e asthma:					
Has the patient had	l a documented trial o	of an inhaled corticost	teroid (ICS) within the	last 120 d	ays? 🛚 Ye	es 🗆 No
Select if the patient	has had a document	ted trial of one of the	following controller me	edications	within the I	last 120 days:
	eta 2 agonist (LABA)					
☐ LABA/ICS con		- (LANAA)				
☐ Long-acting m	iuscarinic antagonist iodifiers	S (LAIVIA)				
☐ Theophylline						
	cribed by or in consu	Itation with an allergis	t/immunologist or gas	stroenterol	naiet2 🗖 V	/os □ No



Dupixent® Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Eosinophilic esophagitis:
Has the patient had a documented trial of a high-dose proton pump inhibitor for at least 8 weeks or swallowed topical steroid (e.g., fluticasone propionate or oral budesonide suspension)? ☐ Yes ☐ No
Was Dupixent prescribed by or in consultation with an allergist/immunologist, pulmonologist, or otolaryngologist?
□ Yes □ No
Eosinophilic esophagitis
Has the patient had a documented trial of a topical corticosteroids or antihistamines within the last 120 days? 🗖 Yes 🗖 No
Was Dupixent prescribed by or in consultation with a dermatologist or allergist/immunologist? ☐ Yes ☐ No
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?
Please note: This request may be denied unless all required information is received.



Enbrel® Prior Authorization Request Form (Page 1 of 2)
DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED Member Information (required) Provider Information (required) Member Name: Provider Name: Insurance ID#: NPI#: Specialty: Office Phone: Date of Birth: Street Address: Office Fax: City: Office Street Address: State: Zip: Phone: Citv: State: Zip: Medication Information (required) Dosage Form: Medication Name: Strenath: ☐ Check if requesting brand Directions for Use: ☐ Check if request is for continuation of therapy Clinical Information (required) Select the diagnosis below: □ Active ankylosing spondylitis (AS) □ Active psoriatic arthritis (PsA) ☐ Moderate to severe chronic plaque psoriasis (PsO) ☐ Moderately to severely active polyarticular juvenile idiopathic arthritis (pJIA) ☐ Moderately to severely active rheumatoid arthritis (RA) Other diagnosis: ICD-10 Code(s): Clinical information: Select if the requested medication is prescribed by or in consultation with one of the following specialists: ■ Dermatologist ■ Rheumatologist Will the requested medication be used in combination with another biologic agent or targeted immunomodulator?

Yes
No For active ankylosing spondylitis (AS), also answer the following: Has the patient had an inadequate response to, intolerance to, or contraindication to one or more non-steroidal anti-inflammatory drugs (NSAIDs)? ☐ Yes ☐ No For active psoriatic arthritis (PsA), also answer the following: Has the patient had an inadequate response to, intolerance to, or contraindication to methotrexate?

Yes No For moderate to severe chronic plaque psoriasis (PsO), also answer the following: Has the patient had an inadequate response to, intolerance to, or contraindication to conventional therapy with at least one of the following: phototherapy or one or more oral systemic treatments (i.e., methotrexate, cyclosporine, acitretin, sulfasalazine)? ☐ Yes ☐ No For moderately to severely active polyarticular juvenile idiopathic arthritis (pJIA), also answer the following: Has the patient had an inadequate response to, intolerance to, or contraindication to one or more non-biologic disease modifying antirheumatic drugs (DMARDs)? ☐ Yes ☐ No For moderately to severely active rheumatoid arthritis (RA), also answer the following:



Enbrel® Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Quantity limit requests:
What is the quantity requested per MONTH? _____
What is the reason for exceeding the plan limitations?
□ Titration or loading dose purposes
□ Patient is on a dose-alternating schedule (e.g., one tablet in the morning and two tablets at night, one to two tablets at bedtime)
□ Requested strength/dose is not commercially available
□ Other: _____

Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?

Please note: This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Enspryng® Prior Authorization Request Form
DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)				Provider Information (required)			
Member Name:			Provider Name	:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street A	ddress:			
Phone:	1	I	City:	State:		Zip:	
		Medication	Information (re	auired)			
Medication Name:			Strength:	4	Dosage F	form:	
☐ Check if requesting	brand		Directions for U	Jse:			
☐ Check if request is	for continuation c	of therapy					
		Clinical Ir	nformation (requi	ired)			
Select the diagnos	sis below:						
□ Neuromyelitis op	`	,					
Other diagnosis:			IC	D-10 Code(s): _			
Clinical informatio							
□ Neurologist	Other		consultation with one				
Will the requested n	nedication be use	ed in combination wi	th another biologic ag	jent? □ Yes □	No		
Are there any other com this review?	ments, diagnoses, s	symptoms, medications	tried or failed, and/or an	y other information	the physicia	n feels is important to	
Please note: This r	equest may be denie	ed unless all required infor	mation is received				



Fasenra[™] Prior Authorization Request Form

Mem	ber Inform	ation (required)	Pro	Provider Information (required)				
Member Name:			Provider Name:					
Insurance ID#:			NPI#:		Specialty:			
Date of Birth:			Office Phone:					
Street Address:			Office Fax:					
City:	State:	Zip:	Office Street Ac	ldress:				
Phone:			City:	State:		Zip:		
		Medication I	nformation (req	uired)				
Medication Name:			Strength:	<i>'</i>	Dosage F	orm:		
☐ Check if requestir	ng brand		Directions for U	Directions for Use:				
☐ Check if request i	s for continuatio r	n of therapy						
		Clinical Inf	ormation (requir	ed)				
Select the diagno	sis below:							
□ Severe asthma	with an eosinor	philic phenotype						
□ Other diagnosi	s:		ICD-10) Code(s):				
dose inhaled cortic	perienced inade costeroid (ICS) a act or leukotriene	quate control of asthma and controlled medication receptor antagonist)?	on (long-acting beta2	agonist (LABA)	or high-dos	se LABA/IČS		
Is Fasenra prescri	bed by or in con	sultation with a rheumat	tologist, pulmonologi	st, allergist, or ir	mmunologis	t? LIYes LING		
Are there any other co his review?	mments, diagnose:	s, symptoms, medications t	ried or failed, and/or any	other information	the physicia	n feels is important		
Please note: Th	is request may be d	enied unless all required infor	mation is received.					



Humira® Prior Authorization Request Form (Page 1 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:	:		
Phone:			City:	State:		Zip:
		Madiaatian Info	rmation			·
Medication Name:	L	Medication Info	Strength:		Dosage F	orm:
					Dosage	OIIII.
☐ Check if requesting	g brand for continuation of the	*****	Directions for Use:			
□ Check if request is	101 Continuation of the		nation			
		Clinical Inform	Mation (required)			
Select the diagnosis						
☐ Active ankylosing	•					
☐ Active psoriatic art	ninus (PSA) e chronic plaque psorias	rie .				
		va (e.g., Hurley Stage II o	or III)			
	erely active Crohn's disc	,	or iii)			
1	•	juvenile idiopathic arthri	tis (JIA)			
I	erely active rheumatoid	•	(
I	erely active ulcerative co	, ,				
■ Non-infectious uve	eitis					
Other diagnosis: _			ICD-10 Cod	de(s):		
Clinical information:						
Dermatologist	Gastroentero	•	almologist 🔲 F	Rheumatolo	•	
Will the requested me	dication be used in com	bination with another bid	ologic agent or targeted	immunomo	dulator? 🔲 🕻	Yes 🗆 No
I		o answer the following				
Has the patient had a (NSAIDs)?		to, intolerance to, or con	traindication to one or m	ore non-ste	eroidal anti-ii	nflammatory drugs
For active psoriatic	arthritis (PsA), also an	swer the following:				
	· · · · · · · · · · · · · · · · · · ·	to, intolerance to, or con		exate? 🛚 Y	es 🗆 No	
		oriasis (PsO), also ans				
	py or one or more oral s	to, intolerance to, or con ystemic treatments (i.e.,				st one of the
For moderate to sev	ere hidradenitis suppu	ırativa, also answer the	e following:			
	n inadequate response ectable steroid therapy?	to, intolerance to, or con Lack December 2	traindication to one or m	ore of the fo	ollowing: ora	al or topical antibiotic
For moderately to se	everely active Crohn's	disease, also answer t	he following:			
	n inadequate response	to, intolerance to, or con	traindication to one or m	ore immuno	suppressive	e agents (e.g.,



Humira® Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

For moderately to severely active polyarticular juvenile idiopathic arthritis (pJIA), also answer the following:
Has the patient had an inadequate response to, intolerance to, or contraindication to one or more non-biologic disease modifying anti-rheumatic drugs (DMARDs)? Yes No
For moderately to severely active rheumatoid arthritis (RA), also answer the following:
Has the patient had an inadequate response to, intolerance to, or contraindication to one or more non-biologic disease modifying anti-rheumatic drugs (DMARDs)? Yes No
For moderately to severely active ulcerative colitis, also answer the following:
Has the patient had an inadequate response to, intolerance to, or contraindication to conventional therapy with one or more of the following: corticosteroids (i.e., prednisone, methylprednisolone), 5-ASAs (i.e., mesalamine, sulfasalazine, balsalazide, olsalazine), non-biologic DMARDs (i.e., azathioprine, methotrexate, mercaptopurine)? Yes No
For non-infectious uveitis, also answer the following:
Has the patient had an inadequate response to, intolerance to, or contraindication to one or more of the following: methotrexate, mycophenolate, azathioprine, cyclosporine, tacrolimus, cyclophosphamide? Yes No
Quantity limit requests:
What is the quantity requested per MONTH?
What is the reason for exceeding the plan limitations? ☐ Titration or loading dose purposes ☐ Patient is on a dose-alternating schedule (e.g., one tablet in the morning and two tablets at night, one to two tablets at bedtime) ☐ Requested strength/dose is not commercially available ☐ Other:
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?

This request may be denied unless all required information is received. Please note:



Ilaris® Prior Authorization Request Form

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name	Provider Name:		
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street A	ddress:		
Phone:			City:	State:	Zip:	
		Medication I	nformation (re	aguired)		
Medication Name:		Modroation	Strength:	squireu)	Dosage Form:	
☐ Check if requesting	g brand		Directions for U	Jse:		
	for continuation of th	nerapy	Birodiono for v			
		Clinical Inf	formation (requ			
syndrome (MWS)] Tumor necrosis far deficiency (MKD) Still's disease Other diagnosis: Clinical information Select if the requeste Allergist/Immun	ctor receptor associate or familial mediterrane : d medication is diagno	ed periodic syndrome an fever sed by, or upon cons gist	e or hyperimmunoglob ICD sultation with or recom Rheumatologist	nulin D syndrome (In-10 Code(s):		
Will the requested me	edication be used in co	mbination with anoth	er biologic agent? 📮	Yes 🗆 No		
_				•	steroidal anti-inflammatory drugs	
Are there any other com this review?	nments, diagnoses, sym	ptoms, medications ti	ried or failed, and/or an	ny other information	n the physician feels is important t	

Please note: This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Ilumya[™] Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address			
Phone:		1	City:	State:	Zip:	
		Medication Info	ormation (required)			
Medication Name:			Strength:		Dosage Form:	
☐ Check if requesting			Directions for Use:			
☐ Check if request is f	or continuation of the	rapy				
		Clinical Inforr	nation (required)			
Select the diagnos	is below:					
■ Moderate-to-seven						
☐ Other diagnosis:			ICD-10	Code(s):		
Clinical information						
	•	with a dermatologist?				
•		nother biologic agent		,.		
	ototherapy or one or r	nse to, intolerance to, more oral systemic tre			nal therapy with at least one osporine, acitretin,	
Are there any other con this review?	nments, diagnoses, sym	ptoms, medications tried	or failed, and/or any othe	er information	the physician feels is important to	
Please note: This	request may be denied un	less all required information	n is received.			



Kevzara® Prior Authorization Request Form
DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:	!		
Phone:	<u> </u>	1	City:	State:		Zip:
		Medication Info	rmation (required)			
Medication Name:			Strength:		Dosage Fo	orm:
☐ Check if requesting	brand		Directions for Use:			
☐ Check if request is	for continuation of the	rapy				
		Clinical Inforr	nation (required)			
Select the diagnos Moderately to se	is below: verely active rheuma	toid arthritis (RA)				
Other diagnosis:			ICD-10 Cod	le(s):		
Clinical information	n:					
Is Kevzara prescribe	ed by or in consultation	on with a rheumatologi	st? 🛘 Yes 🗘 No			
Will Kevzara be use	d in combination with	another biologic ager	nt? 🛘 Yes 🗘 No			
	an inadequate respo natic drugs (DMARDs	nse to, intolerance to, s)? □ Yes □ No	or contraindication to	one or mo	re non-biol	ogic disease
Are there any other con this review?	nments, diagnoses, sym	ptoms, medications tried	or failed, and/or any othe	r information	n the physicia	an feels is important to

This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.

Please note:



Kineret® Prior Authorization Request Form

Member Information (required)			Pro	Provider Information (required)			
Member Name:			Provider Name	Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street A	ddress:			
Phone:			City:	State:	Zip:		
		Medication	Information (re	quired)			
Medication Name:			Strength:	· · · · · · · · · · · · · · · · · · ·	Dosage Form:		
☐ Check if requesting	g brand		Directions for U	Jse:			
☐ Check if request is	for continuatio	n of therapy					
		Clinical Ir	nformation (requi	ired)			
neurologist, or other r Will the requested r For moderately to so	leukin-1 recepto by, or upon consideration be reverely active remainded in the remainded in the remainded in the receptor and inadequate remainded in the remainde	sultation with or recommended in combination with the unatoid arthritis (RA sponse to, intolerance to,	endation of, an immunolo ith another biologic ag), also answer the follo	gent? Yes owing:	matologist, rheumatologist, l No ologic disease modifying anti-		
☐ Requested strengt☐ Other: Are there any other com	equested per Mo for exceeding to g dose purposes se-alternating so th/dose is not co	ne plan limitations? hedule (e.g., one tablet ir mmercially available			e to two tablets at bedtime) the physician feels is important to		
this review?							

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262.



Nucala® Prior Authorization Request Form

	ber Informa			Provider Information (required)			
Member Name:			Provider Name:				
Insurance ID#:			NPI#:	NPI#: Specialty:			
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Addre	Office Street Address:			
Phone:			City:	State:	Zip:		
		Modioation	Information				
Medication Name:		Medication	Information (require Strength:	ed)	Dooggo Form:		
					Dosage Form:		
☐ Check if requesting	-	- £ 41	Directions for Use:				
☐ Check if request is	for continuation (41				
		Clinical In	formation (required)				
Select the diagnos							
□ Severe asthma	with an eosinoph	ilic phenotype					
Eosinophilic gra	ınulomatosis with	polyangiitis (Churg-S	Strauss Syndrome)				
Hypereosinophi	lic syndrome						
Chronic rhinosir	nusitis with nasal	polys (CRSwNP)					
Other diagnosis	i:		ICD-10 C	ode(s):			
Clinical information							
Is Nucala prescribe	ed by or in consul	tation with a rheumate	ologist, pulmonologist, a	llergist, or im	nmunologist? 🛚 Yes 🗘 No		
For severe asthma	a with an eosino	philic phenotype, a	so answer the following	ng:			
		uate control of asthmaedication? ☐ Yes ☐		inimum of thi	ree months use of a high		
Has the patient had months? Yes		nma exacerbations re	quiring medical interven	tion within th	ne past 12		
For chronic rhinos	sinusitis with na	sal polyps (CRSwN	P), also answer the fol	lowing:			
Has the patient exp	perienced inadeq	uate response to nasa	al corticosteroids? 🗖 Y	es 🛭 No			
are there any other comminis review?	nents, diagnoses, s	ymptoms, medications tr	ied or failed, and/or any oth	er information	the physician feels is important to		
Please note: This i	request may be denie	ed unless all required inform	nation is received				

For urgent or expedited requests please call 1-855-401-4262.



Olumiant® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)				
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Address:	:			
Phone:			City:	State:		Zip:	
		Medication Info	rmation (required)				
Medication Name:			Strength:		Dosage Fo	orm:	
☐ Check if requesting	brand		Directions for Use:				
☐ Check if request is f	for continuation of the	rapy					
		Clinical Inforr	nation (required)				
Select the diagnos	is below:						
Moderately to se	verely active rheuma	toid arthritis (RA)					
Other diagnosis:			ICD-10 Cod	ICD-10 Code(s):			
Clinical information	n:						
Is Olumiant prescrib	ed by or in consultati	on with a rheumatolog	jist? 🛘 Yes 🗘 No				
Will Olumiant be use	ed in combination with	h another biologic age	nt? 🛘 Yes 🗘 No				
Has the patient had	an inadequate respo	nse to, intolerance to,	or contraindication to	methotrex	ate? 🛚 Ye	s 🗆 No	
Are there any other con this review?	nments, diagnoses, sym	ptoms, medications tried	or failed, and/or any othe	r information	the physicia	an feels is important to	
Please note: This	request may be depied up	place all required information	a in received				

Please note:

This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262.



Orencia® Prior Authorization Request Form (Page 1 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

	DO NOT COPY FOR	FUTURE USE. FORMS A	RE UPDATED FREQUE	ENILY AND MAY BE	BARCODED		
Member Information (required)			Pro	Provider Information (required)			
Member Name:			Provider Name	Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Ad	ddress:			
Phone:			City:	State:	Zip:		
		Medication I	nformation (re	quired)			
Medication Name:		modioation	Strength:	quireu)	Dosage Form:		
☐ Check if requesting	brand		Directions for U	Jse:			
☐ Check if request is		therapy					
		Clinical Inf	ormation (requi	ired)			
Select the diagnosis	s below:		` ·	•			
☐ Active psoriatic art							
■ Moderately to seven	erely active polyartic	cular juvenile idiopathic a	arthritis (pJIA)				
■ Moderately to seven		toid arthritis (RA)					
☐ Other diagnosis: _			ICD-	·10 Code(s):			
Clinical information							
<u> </u>	d medication is pres	cribed by or in consultat	ion with one of the fol	llowing specialists:			
DermatologistRheumatologist	•						
		combination with another	er biologic agent?	Yes □ No			
-		o answer the following					
Has the patient had a	ın inadequate respor	nse to, intolerance to, or	contraindication to m	ethotrexate? 🗖 Y	es 🗆 No		
For moderately to se	everely active poly	articular juvenile idiop	athic arthritis (pJIA)	, also answer the	following:		
Has the patient had a rheumatic drugs (DM.			contraindication to or	ne or more non-bio	logic disease modifying anti-		
		matoid arthritis (RA), a					
Has the patient had a rheumatic drugs (DM.			contraindication to or	ne or more non-bio	logic disease modifying anti-		
Quantity limit reque What is the quantity re		-LIO					
What is the reason f	•						
☐ Titration or loading	•						
Patient is on a dos	se-alternating sched		e morning and two ta	blets at night, one	to two tablets at bedtime)		
☐ Requested strengt	th/dose is not comm	ercially available					
Other:							



Orencia® Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Are there any of this review?	other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important
Please note:	This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.



Otezla® Prior Authorization Request Form

DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:	:		
Phone:	<u> </u>	. L	City:	State:	Zip:	
		Medication Info	rmation (required)			
Medication Name:			Strength:		Dosage Form:	
☐ Check if requesting	brand		Directions for Use:			
☐ Check if request is	for continuation of the	erapy				
		Clinical Inform	nation (required)			
Select the diagnosis	below:					
Active psoriatic artl	hritis (PsA)					
Moderate to severe	e chronic plaque psoria	sis (PsO)				
Other diagnosis:			ICD-10 Cod	de(s):		
Clinical information:						
Select if the requested Dermatologist	I medication is prescrib ☐ Rheumatologis	ed by or in consultation vet	with one of the following	specialists:		
Will the requested me	dication be used in com	nbination with another bid	ologic agent? 🛚 Yes 🛚	l No		
-	arthritis (PsA), also an					
·		contraindication, or into		? 🗆 Yes 🗆	l No	
For moderate to seve	ere plaque psoriasis (PsO), also answer the f	following:			
		contraindication, or intoleatments (i.e., methotrex			at least one of the following: nzine)?	
Are there any other corthis review?	nments, diagnoses, sym	ptoms, medications tried	or failed, and/or any othe	er information	n the physician feels is important to	
Places note: This	request may be denied up	place all required information	o in roppiyed			

<u>Please note</u>: This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Rinvoq® Prior Authorization Request Form

Insurance ID#: Date of Birth: Street Address: City: State: Zip: Phone: Medication Info Medication Name: Check if requesting brand Check if request is for continuation of therapy Clinical Inform Select the diagnosis below: Moderately to severely active rheumatoid arthritis (RA) Moderately to severely active ulcerative colitis Active psoriatic arthritis Active ankylosing spondylitis Active atopic dermatitis Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w Dermatologist Gastroenterologist Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and altas the patient had an inadequate response to, intolerance to, or contribumira, Simponi, Remicade, etc)?	Strength: Directions for Use:	State:	Specialty: Zip: Dosage Form:
Date of Birth: Street Address: City: State: Zip: Phone: Medication Info Medication Name: Check if requesting brand Check if request is for continuation of therapy Clinical Inform Select the diagnosis below: Moderately to severely active rheumatoid arthritis (RA) Moderately to severely active ulcerative colitis Active psoriatic arthritis Active ankylosing spondylitis Active atopic dermatitis Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w Dermatologist Gastroenterologist Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex) For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a thas the patient had an inadequate response to, intolerance to, or contri	Office Phone: Office Fax: Office Street Address: City: City: City	State:	Zip:
Street Address: City: State: Zip: Phone: Medication Info Medication Name: Check if requesting brand Check if request is for continuation of therapy Clinical Inform Select the diagnosis below: Moderately to severely active rheumatoid arthritis (RA) Moderately to severely active ulcerative colitis Active psoriatic arthritis Active ankylosing spondylitis Active atopic dermatitis Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w	Office Fax: Office Street Address: City: Ci	State:	,
City: State: Zip: Phone: Medication Info Medication Name: Check if requesting brand Check if request is for continuation of therapy Clinical Inform Select the diagnosis below: Moderately to severely active rheumatoid arthritis (RA) Moderately to severely active ulcerative colitis Active psoriatic arthritis Active ankylosing spondylitis Active atopic dermatitis Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Clinical information: Select if the requested medication is prescribed by or in consultation w Cli	Office Street Address: City: Tmation (required) Strength: Directions for Use:	State:	,
Phone: Medication Info Medication Name: Check if requesting brand Check if request is for continuation of therapy Clinical Inform Select the diagnosis below: Moderately to severely active rheumatoid arthritis (RA) Moderately to severely active ulcerative colitis Active psoriatic arthritis Active ankylosing spondylitis Active ankylosing spondylitis Active atopic dermatitis Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w Dermatologist Gastroenterologist Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and alter the patient had an inadequate response to, intolerance to, or contri	City: Tmation (required) Strength: Directions for Use:	State:	,
Medication Name: Check if requesting brand Check if request is for continuation of therapy Clinical Inform Select the diagnosis below: Moderately to severely active rheumatoid arthritis (RA) Moderately to severely active ulcerative colitis Active psoriatic arthritis Active ankylosing spondylitis Active atopic dermatitis Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w Dermatologist Gastroenterologist Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrey For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a thas the patient had an inadequate response to, intolerance to, or contri	rmation (required) Strength: Directions for Use:		,
Medication Name: ☐ Check if requesting brand ☐ Check if request is for continuation of therapy Clinical Inform Select the diagnosis below: ☐ Moderately to severely active rheumatoid arthritis (RA) ☐ Moderately to severely active ulcerative colitis ☐ Active psoriatic arthritis ☐ Active ankylosing spondylitis ☐ Active atopic dermatitis ☐ Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w ☐ Dermatologist ☐ Gastroenterologist ☐ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a thas the patient had an inadequate response to, intolerance to, or contra	rmation (required) Strength: Directions for Use:		,
Medication Name: ☐ Check if requesting brand ☐ Check if request is for continuation of therapy Clinical Inform Select the diagnosis below: ☐ Moderately to severely active rheumatoid arthritis (RA) ☐ Moderately to severely active ulcerative colitis ☐ Active psoriatic arthritis ☐ Active ankylosing spondylitis ☐ Active atopic dermatitis ☐ Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w ☐ Dermatologist ☐ Gastroenterologist ☐ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a thas the patient had an inadequate response to, intolerance to, or contra	Strength: Directions for Use:		Dosage Form:
□ Check if requesting brand □ Check if request is for continuation of therapy Clinical Inform Select the diagnosis below: □ Moderately to severely active rheumatoid arthritis (RA) □ Moderately to severely active ulcerative colitis □ Active psoriatic arthritis □ Active ankylosing spondylitis □ Active atopic dermatitis □ Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w □ Dermatologist □ Gastroenterologist □ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrey For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a thas the patient had an inadequate response to, intolerance to, or contractions.	Directions for Use:		Dosage Form:
☐ Check if request is for continuation of therapy Clinical Inform Select the diagnosis below: ☐ Moderately to severely active rheumatoid arthritis (RA) ☐ Moderately to severely active ulcerative colitis ☐ Active psoriatic arthritis ☐ Active ankylosing spondylitis ☐ Active atopic dermatitis ☐ Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w ☐ Dermatologist ☐ Gastroenterologist ☐ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a thas the patient had an inadequate response to, intolerance to, or contri			
Select the diagnosis below: Moderately to severely active rheumatoid arthritis (RA) Moderately to severely active ulcerative colitis Active psoriatic arthritis Active ankylosing spondylitis Active atopic dermatitis Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w Dermatologist Gastroenterologist Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrey) For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a thas the patient had an inadequate response to, intolerance to, or contri	ation (required)		
Select the diagnosis below: ☐ Moderately to severely active rheumatoid arthritis (RA) ☐ Moderately to severely active ulcerative colitis ☐ Active psoriatic arthritis ☐ Active ankylosing spondylitis ☐ Active atopic dermatitis ☐ Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w ☐ Dermatologist ☐ Gastroenterologist ☐ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex) For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a thas the patient had an inadequate response to, intolerance to, or contra	ation (required)		
Select the diagnosis below: ☐ Moderately to severely active rheumatoid arthritis (RA) ☐ Moderately to severely active ulcerative colitis ☐ Active psoriatic arthritis ☐ Active ankylosing spondylitis ☐ Active atopic dermatitis ☐ Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w ☐ Dermatologist ☐ Gastroenterologist ☐ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex) For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a thas the patient had an inadequate response to, intolerance to, or contractions.	(
□ Moderately to severely active rheumatoid arthritis (RA) □ Moderately to severely active ulcerative colitis □ Active psoriatic arthritis □ Active ankylosing spondylitis □ Active atopic dermatitis □ Other diagnosis: □ Clinical information: Select if the requested medication is prescribed by or in consultation w □ Dermatologist □ Gastroenterologist □ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex) For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a thas the patient had an inadequate response to, intolerance to, or contri		D 40 0 . I. ()	
□ Moderately to severely active ulcerative colitis □ Active psoriatic arthritis □ Active ankylosing spondylitis □ Active atopic dermatitis □ Other diagnosis: □ Clinical information: Select if the requested medication is prescribed by or in consultation w □ Dermatologist □ Gastroenterologist □ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a Has the patient had an inadequate response to, intolerance to, or contr		D 40 0 . I. ()	
□ Active psoriatic arthritis □ Active ankylosing spondylitis □ Active atopic dermatitis □ Other diagnosis: □ Clinical information: Select if the requested medication is prescribed by or in consultation w □ Dermatologist □ Gastroenterologist □ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a Has the patient had an inadequate response to, intolerance to, or contr		D 40 O . I . ()	
□ Active atopic dermatitis □ Other diagnosis: □ Clinical information: Select if the requested medication is prescribed by or in consultation w □ Dermatologist □ Gastroenterologist □ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotres) For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a Has the patient had an inadequate response to, intolerance to, or contr		D 40 0 1 ()	
☐ Other diagnosis: Clinical information: Select if the requested medication is prescribed by or in consultation w ☐ Dermatologist ☐ Gastroenterologist ☐ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a Has the patient had an inadequate response to, intolerance to, or contr		D 40 0 1 ()	
Clinical information: Select if the requested medication is prescribed by or in consultation w ☐ Dermatologist ☐ Gastroenterologist ☐ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a Has the patient had an inadequate response to, intolerance to, or contr		D 40 0 1 / \	
Select if the requested medication is prescribed by or in consultation w Dermatologist Gastroenterologist Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a Has the patient had an inadequate response to, intolerance to, or contr	IC	D-10 Code(s):	
☐ Dermatologist ☐ Gastroenterologist ☐ Rheumatolog Will Rinvoq be used in combination with another biologic agent, Janus potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a Has the patient had an inadequate response to, intolerance to, or contri			
potent immunsuppressants (e.g., azathioprine, cyclosporine, methotrex For rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, and a Has the patient had an inadequate response to, intolerance to, or contr		specialists:	
Has the patient had an inadequate response to, intolerance to, or contr			
	aindication to one or m	ore TNF block	xers (e.g., Cimzia, Enbrel,
For atopic dermatitis also answer the following: Has the patient had an inadequate response to, intolerance to, or contr	aindication to one com-	noro quatomia	drug product for the
treatment of atopic dermatitis (e.g., Adbry, Dupixent, etc)?		ore systemic	arug product for the
are there any other comments, diagnoses, symptoms, medications tried or his review?			e physician feels is importa

This request may be denied unless all required information is received. Please note:



Siliq® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)				
Member Name:			Provider Name	e :			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Address:				
Phone:	<u> </u>		City:	State:	Zip:		
		Medication In	formation (re	equired)			
Medication Name:			Strength:		Dosage Form:		
☐ Check if requesting	brand		Directions for U	Jse:			
☐ Check if request is t	for continuation of	therapy					
		Clinical Info	rmation (requ	ired)			
Select the diagnos	is below:						
■ Moderate to seve	ere chronic plaque	psoriasis					
Other diagnosis:			IC	CD-10 Code(s):			
Clinical information	n:						
Is Siliq prescribed by	y or in consultation	n with a dermatologist?	' □ Yes □ No				
Will Siliq be used in	combination with	another biologic agent	? 🗆 Yes 🚨 No				
Has the patient had an inadequate response to, intolerance to, or contraindication to conventional therapy with at least one of the following: phototherapy or one or more oral systemic treatments (i.e., methotrexate, cyclosporine, acitretin, sulfasalazine)? Yes No							
Are there any other conthis review?	nments, diagnoses, s	symptoms, medications tri	ed or failed, and/or a	ny other information	the physician feels is important to		
Please note: This	request may be denie	d unless all required informa	tion is received				



Simponi® Prior Authorization Request Form (Page 1 of 2)

	DO NOT COPY FOR FU	TURE USE. FORMS ARE	UPDATED FREQUENTLY				
Memb	oer Informatio	N (required)	Provider Information (required)				
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Address:				
Phone:			City: State:			Zip:	
		Medication Int	ormation (required)				
Medication Name:		medication in	Strength:)	Dosage F	orm:	
☐ Check if requesting	brand		Directions for Use:				
☐ Check if request is t		erapy	Directions for edg.				
			rmation (required)				
Select the diagnosis	helow:		rriation (required)				
☐ Active ankylosing s							
☐ Active psoriatic arti	•						
☐ Moderately to seve	, ,	l arthritis (RA)					
■ Moderately to seve	•	, ,					
☐ Other diagnosis:	•		ICD-10 Co	de(s):			
Clinical information:							
			n with one of the following umatologist	specialists:			
Will the requested med	dication be used in co	mbination with another I	oiologic agent? 🗖 Yes 🏻	⊒ No			
For active ankylosing	g spondylitis (AS), al	so answer the following	ng:				
Has the patient had ar (NSAIDs)? ☐ Yes ☐		e, contraindication, or in	colerance to one or more	non-steroida	al anti-inflam	matory drugs	
For active psoriatic a	arthritis (PsA), also a	nswer the following:					
Has the patient had ar	n inadequate response	e, contraindication, or int	olerance to methotrexate	? □ Yes □	l No		
I	=		o answer the following:				
Has the patient had ar rheumatic drugs (DMA			olerance to one or more	non-biologic	disease mo	odifying anti-	
l -	-	ive colitis, also answe	_				
			colerance to conventional				
		nisolone), 5-ASAs (i.e., nercaptopurine)? ☐ Ye	mesalamine, sulfasalazin s 🔲 No	e, balsalazio	le, olsalazin	e), non-biologic	
Quantity limit reques What is the quantity re)					
What is the reason for							
☐ Titration or loading							
Patient is on a dose		(e.g., one tablet in the	morning and two tablets a	at night, one	to two tablet	s at	
bedtime)	h/daga ia mataaa	sially available					
☐ Requested strengtl☐ Patient requires a			ırface area [Tonical ann	lications on	lv1		
 Patient requires a greater quantity for the treatment of a larger surface area [Topical applications only] Other:							



Simponi® Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?								
Please note:	This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.							



Skyrizi® Prior Authorization Request Form

Member Information (required)			Provider Information (required)				
Member Name:			Provider Name:				
Insurance ID#:		NPI#:			Specialty	:	
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Address:				
Phone:			City: State: 2		Zip:		
		Medication	Information (req	uired)			
Medication Name:			Strength:	· · · ·	Dosage F	orm:	
☐ Check if reques	ting brand		Directions for Us	se:			
☐ Check if reques	t is for continuatio	n of therapy					
		Clinical In	formation (require	ed)			
Select the diagr	nosis below:						
☐ Moderate to s	severe plaque pso	oriasis					
Active psoriat	ic arthritis						
Moderately to	severely active (Crohn's disease					
☐ Other diagno	sis:		ICI	D-10 Code(s):			
Clinical informa							
Select if the requ		n is prescribed by or in coroenterologist		of the following Other			
Will the requeste	ed medication be	used in combination with	another biologic age	ent? 🗆 Yes 🛭	⊒ No		
		e response to, intolerand one or more oral systemi	1 1 1 11 1	ion to conventi	• •	•	
Are there any other o	comments, diagnose	es, symptoms, medications t	ried or failed, and/or any	other information	n the physicia	an feels is important to	

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262.



Stelara® Prior Authorization Request Form (Page 1 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:	 Phone:		City:	State:	Zip:	
		Medication Info	ormation (
Medication Name:		Medication iiii	Strength:		Dosage Form:	
☐ Check if requesting	n hrand		Directions for Use:		Dodage Form.	
	for continuation of t	herapy	Directions for Ose.			
			mation (
Clinical Information (required) Select the diagnosis below: Active psoriatic arthritis (PsA) Moderate to severe chronic plaque psoriasis Moderately to severely active Crohn's disease Moderately to severely active ulcerative colitis Other diagnosis: ICD-10 Code(s): Clinical information: Select if the requested medication is prescribed by or in consultation with one of the following specialists: Gastroenterologist Rheumatologist Rheumatologist Will the requested medication be used in combination with another biologic agent? Yes No						
-	• •	answer the following: se to. intolerance to. or co	ntraindication to methotr	exate? 🛚 Y	es □ No	
Has the patient had an inadequate response to, intolerance to, or contraindication to methotrexate? No For moderate to severe chronic plaque psoriasis, also answer the following: Has the patient had an inadequate response to, intolerance to, or contraindication to conventional therapy with at least one of the following: phototherapy or one or more oral systemic treatments (i.e., methotrexate, calcipotriene, cyclosporine, acitretin, sulfasalazine, tazarotene, corticosteroid)? No						
_		's disease, also answer	_		thorony (o.g. ozatkiania	
Has the patient had an inadequate response to, intolerance to, or contraindication to one or conventional therapy (e.g., azathioprine, mercaptopurine, methotrexate, corticosteroids)? \(\Q_{\text{Yes}}\) No						
_	•	ntive Colitis, also answe				
Has the patient had an inadequate response to, intolerance to, or contraindication to one or more conventional therapy (e.g., corticosteroids, mesalamine, balsalazide, olsalazine, azathioprine, mercaptopurine, methotrexate)? No						



Stelara® Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

_	it requests: uantity requested per TREATMENT? syringe every weeks
What is the	reason for exceeding the plan limitations? or loading dose purposes
□ Patient is □ Requeste	on a dose-alternating schedule (e.g., one tablet in the morning and two tablets at night, one to two tablets at bedtime) d strength/dose is not commercially available
U Other:	
Are there any of this review?	ther comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to
•	ther comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to
•	ther comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to



Taltz® Prior Authorization Request Form

D	O NOT COPY FOR FUTU	IRE USE. FORMS ARE UP	DATED FREQUENTLY AN	ND MAY BE	BARCODED	
Memb	er Informatior	(required)	Provide	er Infor	mation	(required)
Member Name:			Provider Name:			
Insurance ID#:			NPI#: Specialty:			
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:	hone:		City:	State:		Zip:
		Medication Info	rmation (required)			
Medication Name:			Strength:		Dosage Fo	orm:
☐ Check if requesting			Directions for Use:		<u> </u>	
Check if request is t	for continuation of the	rapy				
		Clinical Inforr	nation (required)			
Select the diagnosis	below:					
Active ankylosing s	spondylitis					
Active psoriatic arth	hritis					
Moderate to severe	e plaque psoriasis					
Non-radiographic a	axial spondyloarthritis w	ith objective of inflamma	tion			
Other diagnosis:			ICD-10 Cod	le(s):		
Clinical information:	l : : :	- 4				
-	medication is prescribe	ed by or in consultation v	vith one of the following	specialists:		
☐ Dermatologist						
☐ Rheumatologist	dication be used in som	bination with another bid	ologio agant? 🗖 Vaa 🖂	l No		
•					owing	
		adiographic axial spon to, intolerance to, or con	·		-	flammatory drugs
(NSAIDs)? \(\text{Ves}\)		to, intolerance to, or con	traindication to one or m	ore non-ste	ioidai aiiti-iii	naminatory drugs
	arthritis, also answer t					
•		to, intolerance to, or con		xate? 🛚 Ye	es 🗆 No	
		also answer the followi	_			
Has the patient had an inadequate response to, intolerance to, or contraindication to conventional therapy with at least one of the following: phototherapy or one or more oral systemic treatments (i.e., methotrexate, cyclosporine, acitretin, sulfasalazine)? Yes No						
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?						

This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262. Please note:



Tremfya® Prior Authorization Request Form OPY FOR FUTURE USE, FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

	er Informatio	n (required)			mation (required)		
Member Name:			Provider Name:				
Insurance ID#:			NPI#:	Specialty:			
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Address:				
Phone:			City:	State:	Zip:		
		Medication Info	rmation (required)				
Medication Name:			Strength:		Dosage Form:		
☐ Check if requesting	brand		Directions for Use:				
☐ Check if request is	for continuation of th	erapy					
		Clinical Inforr	nation (required)				
Select the diagnos	is below:						
■ Moderate to seve	ere plaque psoriasis						
■ Moderate to seve	ere psoriatic arthritis	•					
Other diagnosis:			ICD-10	Code(s): _			
Clinical informatio	n:						
Is Tremfya prescribe	ed by or in consultat	ion with a dermatologis	t? □ Yes □ No				
Will Tremfya be use	d in combination wit	h another biologic age	nt? 🗆 Yes 🔲 No				
Has the patient had an inadequate response to, intolerance to, or contraindication to conventional therapy with at least one of the following: phototherapy or one or more oral systemic treatments (i.e., methotrexate, cyclosporine, acitretin, Sulfasalazine, calcipotriene, tazarotene, corticosteroid)? Yes No							
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?							

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262.



Xeljanz[®] & Xeljanz XR[®] Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:	Provider Name:		
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Ad	ldress:		
Phone:			City:	State:	Zip:	
		Medication I	nformation (red	auired)		
Medication Name:			Strength:	, ,	Dosage Form:	
☐ Check if requesting	brand		Directions for U	se:		
☐ Check if request is	for continuation o	f therapy				
		Clinical Inf	ormation (requir	red)		
☐ Active ankylosing s☐ Other diagnosis: ☐ Clinical information: Select if the requested☐ Dermatologist☐ Gastroenterolog☐ Rheumatologist☐ Will the requested me Has the patient had all other contents.	hritis erely active rheuma erely active ulcerative erely active polyartic espondylitis d medication is pres ist dication be used in n inadequate respo	ve colitis cular juvenile idiopathic a scribed by or in consultar combination with anothe	ICD- tion with one of the foll er biologic agent?	Yes □ No		
Are there any other com this review?	ments, diagnoses, s	ymptoms, medications tr	ied or failed, and/or any	other information	n the physician feels is important to	

<u>Please note</u>: This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Xolair® Prior Authorization Request Form

		TURE USE. FORMS ARE I			
Member Information (required)		Provider Information (required)			
Member Name:			Provider Name:		
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone:		
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street Address:		
Phone:			City:	State:	Zip:
		Medication Info	ormation (required)		
Medication Name:			Strength:		Dosage Form:
☐ Check if requesting	brand		Directions for Use:		
☐ Check if request is	for continuation of th	erapy			
		Clinical Infor	mation (required)		
Other diagnosis: For asthma, answer Does the patient have	the following: a positive skin test or	in vitro reactivity to a per	ICD-10 Code		
Are the patient's symp	otoms inadequately cor	ntrolled with inhaled cortion			
For chronic idiopath Does the patient rema Is Xolair prescribed by Quantity limit reques	ic urticaria, answer the symptomatic despite or in consultation with sts:	ne following: e H1 antihistamine treatn a dermatologist, rheuma	nent?		immunologist? ☐ Yes ☐ No
What is the reason fo ☐ Titration or loading ☐ Patient is on a dos		limitations? (e.g., one tablet in the m	orning and two tablets a	t night, one t	to two tablets at bedtime)
	mments, diagnoses, syn	nptoms, medications tried	or failed, and/or any othe	er information	n the physician feels is important to
Please note: This	request may be denied u	nless all required information	n is received.		



Juxtapid® Prior Authorization Request Form

Member Information (required)			Provider Information (required)				
Member Name:		Provider Name:					
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:		1		
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Address:				
Phone:		I	City:	State:	Zip:		
		Medication	Information (re	equired)			
Medication Name:			Strength:		Dosage Form:		
☐ Check if requestin	g brand		Directions for U	se:			
☐ Check if request is	for continuation	on of therapy					
		Clinical I	nformation (requ	uired)			
Is the requested m Has the patient had	eline LDL-C le edication pres d trial and failu	evel greater than or equ cribed by or in consulta re of Praluent or Repat d failure with Praluent o	ition with a cardiologis ha? □ Yes □ No	st or endocrinolo			
What is the medica	ıl rationale for	use of Juxtapid over Pr	aluent or Repatha? _				
are there any other con	nments, diagnos	es, symptoms, medications	s tried or failed, and/or ar	ny other informatio	on the physician feels is important		

This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262. Please note:



Extina, XolgelTM & XolegelTM Duo Prior Authorization Request Form

Member Information (required)		Provider Information (required)				
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Addr	ress:		
Phone:			City:	State:		Zip:
		Medication In	formation (requ	uired)		
Medication Name:			Strength:	,	Dosage F	orm:
☐ Check if requesting	g brand		Directions for Use):		
☐ Check if request is	for continuation of	of therapy				
		Clinical Info	ormation (require	ed)		
Select the diagn	osis below:					
		inocompetent patients				
Other diagnos	is:		ICD-10 Code	(s):		
Clinical informat	_					
Has the patient had 120 days?		ilure (a minimum of 60	day trial) of ketoc	onazole crea	ım or sham	npoo in the past
Quantity limit re						
		er MONTH?				
		ng the plan limitation				
•	es a larger quan	tity to cover a larger su	urface area			
Other:						
Are there any other co this review?	mments, diagnoses	, symptoms, medications trie	ed or failed, and/or any	other informatio	n the physici	an feels is important to
Please note: This	s request may be den	ied unless all required informat	ion is received.			

For urgent or expedited requests please call 1-855-401-4262.



Topical onychomycosis agents Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)		Pr	Provider Information (required)					
Member Name:			Provider Name:					
Insurance ID#:		NPI#:		Specialty:				
Date of Birth:			Office Phone:					
Street Address:			Office Fax:					
City:	State:	Zip:	Office Street Address:					
Phone:			City:	State:		Zip:		
		Medication In	formation (re	equired)				
Medication Name:			Strength:		Dosage F	orm:		
☐ Check if requesti	•		Directions for U	Directions for Use:				
☐ Check if request	is for continuation of	therapy						
		Clinical Info	rmation (requ	ired)				
Select the diag	nosis below:							
☐ Onychomyco	sis of the toenails							
Other diagno	sis:		ICD-10 Code(s):					
Clinical informa	ation:							
Has the patient 12 months?		ure of 90 days of terbi	nafine tablets ar	nd 90 days of to	pical cicl	opirox in the last		
Are there any other of this review?	comments, diagnoses, s	symptoms, medications tried	d or failed, and/or an	ny other information	the physici	an feels is important to		
		d unless all required informati						



Luzu® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)					
Member Name:			Provider Na	me:				
Insurance ID#:			NPI#:		Specialty:			
Date of Birth:			Office Phone	e:				
Street Address:	Street Address:							
City:	State:	Zip:	Office Street	Office Street Address:				
Phone:			City:	State:	Zip:			
		Medication I	nformation	(required)				
Medication Name:			Strength:		Dosage Form:			
☐ Check if requesting	•		Directions fo	Directions for Use:				
☐ Check if request is	for continuation	on of therapy						
		Clinical Inf	ormation (r	equired)				
What is the patie	ent's diagno	sis for the medication	being request	ed? (Mandatory	()			
ICD-10 Code(s)	[Mandatory]:						
Medication histo	ory:							
Has the patient tr	ied and faile	d two topical antifungal a	agents in the la	st 365 days? 🗖	Yes □ No			
Has the patient tr	ied and faile	d two oral antifungal age	ents in the last	365 days? □ Ye s	s 🗖 No			
Are there any other co this review?	mments, diagno	ses, symptoms, medications to	ried or failed, and/o	r any other informatio	on the physician feels is important to			
Please note: This	s request may be	denied unless all required inform	ation is received.					

For urgent or expedited requests please call 1-855-401-4262.



Oravig® Prior Authorization Request Form

	er Informatio	n (required)		der Info		
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:		L	City:	State:		Zip:
		Medication Inf	ormation (require	d)		
Medication Name:			Strength:		Dosage Fo	orm:
☐ Check if requesting	brand		Directions for Use:			
☐ Check if request is f	or continuation of th	nerapy				
		Clinical Infor	mation (required)			
Select the diagno	sis below:					
□ Local treatment	t of oropharyngea	l candidiasis (OPC)				
Other diagnosis	S:		_ ICD-10 Code(s):			
Clinical informati	on:					
		e of clotrimazole trock	nes, fluconazole tal	olets/suspe	ension, or	nystatin
suspension within		? LI Yes LI No				
Quantity limit req What is the quantit		DAY?				
•	• •	the plan limitations	?			
☐ Titration or load			•			
☐ Patient is on a	dose-alternating s	schedule (e.g., one ta	blet in the morning	and two ta	blets at ni	ght, one to two
tablets at bedtir	,		_			
•	•	commercially available	9			
Other:						
Are there any other com this review?	nments, diagnoses, sy	mptoms, medications tried	or failed, and/or any oth	er information	the physicia	nn feels is important to
Please note: This	request may be denied	unless all required information	n is received			



Vusion® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Memb	er Informati	On (required)			rmation (required)		
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Ad	ddress:			
Phone:			City:	State:	Zip:		
		Medication In	formation (r	equired)			
Medication Name:			Strength:		Dosage Form:		
☐ Check if requesting			Directions for U	Jse:			
☐ Check if request is	for continuation of		_				
		Clinical Info	rmation (requ	uired)			
Select the diagno							
•	•	dermatitis complicated	•				
Other diagnosi			ICD-10 Code(s):				
Clinical informat Has the patient hat the last 30 days?	ad a trial and failu	re (a minimum of 14 o	day trial) to topi	ical nystatin or t	topical OTC miconazole in		
Quantity limit red What is the quant	•	MONTH?					
•		the plan limitations	:7				
		y to cover a larger su					
Other:	• .						
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?							
Place note: This	request may be denied	duplose all required information	on is received				

Please note:

For urgent or expedited requests please call 1-855-401-4262.



Makena® SubQ Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)		Pi	Provider Information (required)				
Member Name	e:		Provider Name:				
Insurance ID#	:		NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address	s:		Office Fax:				
City:	State:	Zip:	Office Street A	Address:			
Phone:			City:	State:	Zip:		
		Medication	Information	(required)			
Medication Na	ime:		Strength:		Dosage Form:		
	questing brand		Directions for	Use:			
☐ Check if red	quest is for continuatio	n of therapy					
		Clinical Ir	nformation (red	quired)			
Select the di	iagnosis below:						
□ Pregnanc	y indication, preterm	birth					
☐ Other diag	gnosis:		ICE	D-10 Code(s):			
 Is the pati Is the ther 	patient have a history ient having a singleto rapy starting between py be continued until	n pregnancy? Q Yes 16 weeks, 0 days and	■ No 20 weeks, 6 days	of gestation?	erm birth(s)? Yes No Yes No ry, which ever occurs first?		
Are there any oth	her comments, diagnose	s, symptoms, medications	tried or failed, and/or	any other information	on the physician feels is important to		
Please note:	This request may be de	nied unless all required infor	mation is received.				



Metozolv® ODT (metoclopramide orally disintegrating tablet [ODT]) **Prior Authorization Request Form**

	DO NOT COPY FOR FUT	URE USE. FORMS ARE U	PDATED FREQUENTLY	AND MAY BE	BARCODED		
Memb	er Information	(required)	Provi	ider Info	rmation (required)		
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Addres	s:			
Phone:		,	City:	State:	Zip:		
		Medication Info	ormation (require	ed)			
Medication Name:			Strength:		Dosage Form:		
☐ Check if requesting	brand		Directions for Use:		L		
☐ Check if request is f	or continuation of the	гару					
		Clinical Infor	mation (required)				
☐ Symptomatic ga	paresis (diabetic ga astroesophageal re		_ ICD-10 Code(s)	:			
Clinical informati	on:						
	d a 30-day trial and	failure of Brand Re	glan or generic m	etocloprami	ide tablet or solution within		
Quantity limit req What is the quantit	y requested per DA						
		ne plan limitations	?				
tablets at bedtir	dose-alternating sc ne)			ງ and two ta	ablets at night, one to two		
Other:	-	-					
Are there any other com this review?	nments, diagnoses, symp	otoms, medications tried	or failed, and/or any otl	ner information	n the physician feels is important to		
Please note: This	request may be denied up	less all required information	n is received				

For urgent or expedited requests please call 1-855-401-4262.



Multiple Sclerosis Prior Authorization Request Form (Page 1 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

	per Information (re				mation	(required)
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State: Z	ip:	Office Street Address:			
Phone:			City:	State:		Zip:
	Me	dication Info	ormation (required)			
Medication Name:			Strength:		Dosage F	orm:
☐ Check if requesting	brand		Directions for Use:			
☐ Check if request is	for continuation of therap	у	-			
Clinical Information (required)						
Select the medication						
□ Ampyra	☐ Betaseron	☐ Gilenya	■ Maver	nclad	□ Re	ebif
☐ Aubagio	□ Copaxone	☐ Glatirame	r □ Mayze	ent	☐ Ta	scenso ODT
□ Avonex	Dalfampridine ER	Glatopa	☐ Plegri		☐ Te	ecfidera
■ Bafiertam	Extavia	Kesimpta	☐ Ponvo	ry		ımerity
					☐ Ze	posia
Select the diagnosis						
Multiple sclerosis						
			ICD-10 Code	e(s):		
Prescriber's special	-					
□ Neurologist	d medication is prescribed I byra (dalfampridine ER) only		with one of the following	specialists:		
For Ampyra (dalfam	pridine ER), also answer	he following:				
Does the patient have	e a history of seizures? 🗖 🗅	'es □ No				
For Aubagio, Avone Plegridy, Ponvory, R	x, Bafiertam, Betaseron, C Rebif, Tecfidera, or Vumer	Copaxone, Extavia, ity, also answer the	Gilenya, Glatiramer, G following:	latopa, Kes	impta, Lem	trada, Mayzent,
Does the patient have a relapsing form of multiple sclerosis, including clinically isolated syndrome, relapsing-remitting disease, or active secondary progressive disease? Yes No						
For mitoxantrone, al	so answer the following:					
· ·	Itiple sclerosis that applies	to the patient:				
Progressive relation	psing multiple sclerosis	•				
	ressive multiple sclerosis					
Worsening relap	Worsening relapsing-remitting multiple sclerosis					



Multiple Sclerosis Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

sclerosis, including relapsing-remitting	disease or active secondary progressive
nended lifetime limit of 2 treatment cour	ses (or 4 treatment cycles total) of
e sclerosis the patient has failed after a	trial of at least 4 weeks, has a contraindication
☐ Gilenya (fingolimod)	Rebif (interferon beta-1a)
☐ Kesimpta (ofatumumab)	Tecfidera (dimethyl fumarate)
■ Lemtrada (alemtuzumab)	☐ Tysabri (natalizumab)
Mayzent (siponimod)	Vumerity (diroximel)
Ocrevus (ocrelizumab)	Zeposia (ozanimod)
☐ Plegridy (peginterferon beta-1a)	
ations? one tablet in the morning and two table available	ets at night, one to two tablets at bedtime)
medications tried or failed, and/or any ot	her information the physician feels is important to
	e sclerosis the patient has failed after a Gilenya (fingolimod) Kesimpta (ofatumumab) Lemtrada (alemtuzumab) Mayzent (siponimod) Ocrevus (ocrelizumab) Plegridy (peginterferon beta-1a) ations? one tablet in the morning and two table available

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262.



Tysabri® Prior Authorization Request Form

	DO NOT COPY FOR	FUTURE USE. FORMS A	RE UPDATED FREQUEN	TLY AND MAY BE I	BARCODED		
Member Information (required)			Pro	Provider Information (required)			
Member Name:			Provider Name:	Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Ad	Office Street Address:			
Phone:		<u> </u>	City:	State:	Zip:		
		Medication	Information (req	ivod\			
Medication Name	:	Medication	Strength:	uirea)	Dosage Form:		
☐ Check if reques	sting brand		Directions for Us	se:			
•	st is for continuation o	f therapy					
		Clinical In	formation (require	ed)			
Crohn's Disea Other Prescriber's spe Select if the reque Neurologist Gastroenterold Other Quantity limit rewhat is the quant What is the reas Titration or load Patient is on a Requested stree	cialty: ested medication is prespected medication is prespected medication is prespected per MON confor exceeding the plant ding dose purposes dose-alternating schedength/dose is not communication.	scribed by or in consulta TH? plan limitations? dule (e.g., one tablet in the consultation)	ation with one of the follo	owing specialists:	to two tablets at bedtime)		
		d unless all required inforn quests please call 1-855-40					



Nasal Steroids Prior Authorization Request Form

DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)				Provider Information (required)			
Member Name:				Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:	:	<u> </u>		
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street	Office Street Address:			
Phone:		I	City:	State:	Zip:		
		Medication	Information	(required)			
Medication Name:			Strength:		Dosage Form:		
☐ Check if requesting	brand		Directions for	Use:			
☐ Check if request is	for continuation	on of therapy					
		Clinical Ir	nformation (re	quired)			
 □ Nasal polyps □ Nonallergic (value) □ Perennial aller □ Seasonal aller □ Other diagnos 	gic rhinitis gic rhinitis		ICD-10 Cc	ode(s):			
Medication histo	•	l failura af a gamaria na	and atomaid in the	naat C mantha?			
-		I failure of a generic na	isai steroid in the	pasi o monins?	LI TES LI NO		
Quantity limit re What is the quant What is the reas	ity requested	d per MONTH?eding the plan limitati	 ions?				
□ Titration or loa	iding dose pu	urposes					
		ating schedule (e.g., or	ne tablet in the mo	orning and two to	ablets at night, one to two		
tablets at bedt	,	s not commercially ava	nilable				
Other:	•	•	abro				
Are there any other co this review?	mments, diagno	ses, symptoms, medications	tried or failed, and/or	any other informatio	n the physician feels is important to		
Please note: This	roquest may be	denied unless all required info	rmation is received				

For urgent or expedited requests please call 1-855-401-4262.



Nascobal® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)		Pi		rmation (required)			
Member Name:			Provider Nam	e:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:		Office Fax:					
City:	State:	Zip:	Office Street A	Address:			
Phone:	I	L	City:	Zip:			
		Medication	Information (equired)			
Medication Name:			Strength:	Strength: Dosage Form:			
☐ Check if req	uesting brand		Directions for	Directions for Use:			
☐ Check if req	uest is for continuatio	n of therapy					
		Clinical In	formation (req	uired)			
Has the pati	ent had a trial and	failure of injectable cy	anocobalamin wit	hin the past 6 m	nonths?		
Are there any or this review?	ther comments, diagnos	es, symptoms, medications	tried or failed, and/or	any other informatio	n the physician feels is important to		
Please note:	, ,	enied unless all required infor					



NuplazidTM Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Pı	rovider Info	rmation	(required)
Member Name:			Provider Name	: :		
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street A	ddress:		
Phone:			City:	State:		Zip:
		Medication Inf	ormation (r	required)		
Medication Name:			Strength:	· · · · · · · · · · · · · · · · · · ·	Dosage Fo	orm:
☐ Check if requesting brand			Directions for Use:			
☐ Check if request is	for continuation of th	erapy				
		Clinical Infor	mation (req	uired)		
Select the diagn	osis below:					
Hallucinations	and delusions ass	ociated with Parkinso	on's disease p	sychosis		
Other diagnos	is:		_ ICD-10 Cod	de(s):		
Clinical informat	tion:					
Is Nuplazid presc	ribed by or in cons	ultation with a neurol	ogist or psych	iatrist? 🛚 Yes	□ No	
Are there any other co	mments, diagnoses, syr	nptoms, medications tried	or failed, and/or a	ny other information	n the physicia	an feels is important to
Please note: This	s request may be denied u	ınless all required information	n is received.			

For urgent or expedited requests please call 1-855-401-4262.



NuvessaTM Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)				Provider Info	rmation	(required)	
Member Name:			Provider Nan	Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone	:	-11		
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street	Address:			
Phone:			City:	State:		Zip:	
		Medication Inf	formation	(required)			
Medication Name:			Strength: Dosage Form:		orm:		
☐ Check if requestin	g brand		Directions for Use:				
☐ Check if request is	for continuation c	of therapy					
		Clinical Info	rmation (re	equired)			
Has the patient h	ad a trial and fa	ilure of metronidazole va	aginal gel 0.7	75% within the pa	st 30 days	? □ Yes □ No	
Are there any other co	omments, diagnoses	s, symptoms, medications tried	or failed, and/or	r any other information	n the physicia	an feels is important to	
Please note: Thi	s request may be den	nied unless all required information	on is received.				

For urgent or expedited requests please call 1-855-401-4262.



Hetlioz® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Men		ation (required)			ermation (required)		
Member Name:			Provider Nam	Provider Name:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:	:			
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street	Address:			
Phone:			City:	State:	Zip:		
		Medication I	nformation	(required)			
Medication Name:			Strength:	· /	Dosage Form:		
☐ Check if reques	ting brand		Directions for	Directions for Use:			
☐ Check if reques	t is for continuatio	n of therapy					
		Clinical Inf	formation (re	quired)			
	Sleep-Wake Dis	order n Smith-Magenis syndro	me				
☐ Other diagno	sis:		10	ICD-10 Code(s):			
	ried and failed a			copiclone, temaze	pam, triazolam, zaleplon,		
Are there any other of this review?	comments, diagnose	es, symptoms, medications to	ried or failed, and/or	any other information	on the physician feels is important to		
Please note: The state of the s	nis request may be de	enied unless all required inform	ation is received.				

For urgent or expedited requests please call 1-855-401-4262.



Nuvigil® (armodafinil) and Provigil® (modafinil) Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name	Provider Name:		
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street A	Office Street Address:		
Phone:			City:	City: State: Zi		
		Medication	Information (required)		
Medication Name:			Strength:		Dosage Form:	
☐ Check if requesting	g brand		Directions for U	Jse:		
☐ Check if request is		of therapy				
		Clinical In	formation (req	uired)		
Select the diagn	osis below:					
_		ated with obstructive	sleep apnea/hypo	opnea syndrom	e	
□ Narcolepsy						
☐ Shift work slee	ep disorder					
			ICD-10 Co	de(s):		
Quantity limit red	•					
•		oer DAY?				
		ing the plan limitation	ons?			
☐ Titration or loa			e tablet in the mo	rning and two t	ablets at night, one to two	
tablets at bedt		rig concadio (c.g., cri		ming and two t	abloto at mgm, one to two	
□ Requested str	ength/dose is r	not commercially avai	ilable			
□ Other:		·				
				any other informatio	on the physician feels is important to	

This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262.

Please note:



Sunosi[™] & Wakix[®] Prior Authorization Request Form

		ation (required)			mation (required)	
Member Name:				Provider Name:		
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:		<u> </u>	
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Ad	Office Street Address:		
Phone:	_		City:	State:	Zip:	
		Medication I	nformation (red	quired)		
Medication Name:		modication	Strength:	quirea	Dosage Form:	
☐ Check if requesting	g brand		Directions for U	Jse:		
☐ Check if request is		of therapy				
		Clinical Inf	ormation (requi	ired)		
Select the diagnosis	s below:					
□ Narcolepsy with e	•	sleepiness				
☐ Obstructive sleep	•					
Other diagnosis: _			ICD-1	0 Code(s):		
		ime sleepiness, answer t				
		at least one of the following mine, methylphenidate?		gents: amphetami	ne/dextroamphetamine,	
Quantity limit reque What is the quantity r		V2				
What is the reason f		· · · · · · · · · · · · · · · · · · ·				
☐ Titration or loading	g dose purposes	•				
	se-alternating sch	nedule (e.g., one tablet in the	ne morning and two ta	blets at night, one	to two tablets at	
bedtime) Requested streng	th/dose is not cor	mmercially available				
□ Patient requires a	greater quantity	for the treatment of a large		al applications on	ily]	
Are there any other com this review?	nments, diagnoses	s, symptoms, medications tr	ied or failed, and/or any	y other information	the physician feels is important to	
Diagon noto: This	request may be de-	aind unloss all required inform	ation is nearly ad			

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262.



Xyrem® Prior Authorization Request Form

			RE UPDATED FREQUENTLY			
	er Informati	On (required)		er Infor	mation (required)	
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:		1	
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address	Office Street Address:		
Phone:	Л		City:	State:	Zip:	
		Medication Ir	nformation (required)			
Medication Name:			Strength:		Dosage Form:	
☐ Check if requesting	brand		Directions for Use:		<u> </u>	
☐ Check if request is	for continuation of	therapy				
		Clinical Info	ormation (required)			
Select the diagnosis	below:		(- 1			
■ Narcolepsy with ca	ıtaplexy					
■ Narcolepsy with ex	•	•				
Other diagnosis:			ICD-10 Code	e(s):		
Clinical Information:						
Is the patient enrolled	in the Xyrem Succe	ess Program? Yes	l No			
	-	sleepiness, answer th	_			
Has the patient had a armodafinil, modafinil,	previous trial of at le dextroamphetamin	east one of the following e, methylphenidate? $lacksquare$	standard stimulant agents: Yes 🔲 No	amphetami	ne/dextroamphetamine,	
Quantity limit reques						
What is the quantity re						
What is the reason fo ☐ Titration or loading		lan limitations?				
		ule (e.g., one tablet in the	e morning and two tablets a	t night, one	to two tablets at	
☐ Requested strengtl	h/dose is not comm	ercially available				
□ Patient requires a g□ Other:		the treatment of a larger	surface area [Topical appl	ications on	ly] 	
Are there any other cor this review?	nments, diagnoses,	symptoms, medications tr	ied or failed, and/or any othe	er information	n the physician feels is important to	

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262.



Onfi® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)		F	Provider Information (required)			
Member Name:		Provider Nam	ne:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone	:		
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street	Address:		
Phone:	1		City:	State:	Zip:	
		Medication I	nformation	(required)		
Medication Name:			Strength:	` '	Dosage Form:	
☐ Check if requesting brand			Directions for	Use:		
□ Check if request	t is for continuatio	n of therapy				
		Clinical Inf	ormation (re	equired)		
☐ Seizures as:	reatment-resista sociated with Le	ant seizure disorder ennox-Gastaut syndrom IO		:		
Prescriber spe	ecialty:					
Is Onfi prescribe	ed by or in cons	sultation with a neurolog	ist? 🛭 Yes 🗖	No		
Are there any other this review?	comments, diagnos	ses, symptoms, medications tri	ied or failed, and/or	any other informatio	n the physician feels is important to	
Please note:	Γhis request may be α	denied unless all required informa	ation is received.			

For urgent or expedited requests please call 1-855-401-4262.

South Dakota
Department of
Social Services

Fax to 1-844-403-1029 Mon-Sat: 7am to 7pm Central

Bepreve[®], Lastacaft[®], Pataday[®], Patanol[®], Pazeo[®] Prior Authorization Request Form

	O NOT COPY FOR	R FUTURE USE. FORMS ARE	UPDATED FREQUEN	TLY AND MAY B	E BARCODED	
Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:	<u>I</u>		City:	State:	Zip:	
		Medication In	formation (red	quired)		
Medication Name:			Strength:	·	Dosage Form:	
☐ Check if requesting	brand		Directions for Us	e:		
☐ Check if request is	for continuation	of therapy				
		Clinical Info	ormation (requir	red)		
Select the diagnos Allergic conjunct Other diagnosis:	ivitis		ICD-10	Code(s):		
Medication history Has the patient had 120 days? ☐ Yes ☐	a 5 day trial of	azelastine, emedastine,	epinastine, generic	olopatadine, c	or ketotifen in the last	
☐ Titration or loadii☐ Patient is on a do bedtime)	requested per of for exceeding ong dose purpos ose-alternating	the plan limitations?	et in the morning an	nd two tablets a	at night, one to two tablets at	
Are there any other combined this review?	ments, diagnoses	symptoms, medications trie	d or failed, and/or any	other informatio	on the physician feels is important to	

<u>Please note</u>: This request may be denied unless all required information is received.



Opzelura[™] Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)		Pro	Provider Information (required)			
Member Name:			Provider Name:	Provider Name:		
Insurance ID#: NPI#:			S	Specialty:		
Date of Birth:	Date of Birth: Office Phone:					
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Ad	Office Street Address:		
Phone:			City:	State:	Zip:	
		Medication	Information (re	equired)		
Medication Name:			Strength:		Oosage Form:	
☐ Check if requesting	g brand		Directions for U	se:		
☐ Check if request is	for continuatio	n of therapy				
ı		Clinical Ir	nformation (requ	ired)		
Select the diagnos	sis below:			· · · · · · · · · · · · · · · · · · ·		
☐ Actopic dermati						
Other diagnosis	:		ICD-	10 Code(s):		
Clinical information	on:					
1. Does the patien	t have greater	than or equal to 3% bo	dy surface area invol	vement? 🔲 Yes	□ No	
		areas (e.g., face, hands				
	•		py with one of the fol	llowing: corticoster	oids, pimecrolimus and/or	
tacrolimus, crisa						
				nase inhibitors, or p	ootent immunosuppressants	
		porine? Tyes No	0			
5. What is the requ	uested quantity	/?			 	
6. How long Will the	e patient be us	sing Opzelura?				
	nments, diagnose	es, symptoms, medications	tried or failed, and/or ar	ny other information th	he physician feels is important to	
this review?						
Please note: This	request may be de	enied unless all required infor	mation is received.			

For urgent or expedited requests please call 1-855-401-4262.



Oracea®, Seysara®, and Solodyn® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Memb	er Information	(required)	Provid	ler Info	rmation	(required)
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:		L	
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:	l		City:	State:		Zip:
		Medication Inf	ormation (required)		
Medication Name:			Strength:		Dosage Fo	orm:
☐ Check if requesting			Directions for Use:			
☐ Check if request is t	for continuation of the					
		Clinical Infor	mation (required)			
☐ Inflammatory les	ions of non-nodular m ions (papules and pus	stules) of rosacea [Or	ne vulgaris [Seysara a acea only] ICD-10 Co	•		
Clinical information				uo(o)		
Has the patient had	a trial and failure (a n		al) of doxycycline mon- lays? ☐ Yes ☐ No	ohydrate, d	loxycycline	hyclate,
minocycline immediate-release, or tetracycline in the last 180 days? ☐ Yes ☐ No Quantity limit requests: What is the quantity requested per DAY? What is the reason for exceeding the plan limitations? ☐ Titration or loading dose purposes ☐ Patient is on a dose-alternating schedule (e.g., one tablet in the morning and two tablets at night, one to two tablets at bedtime) ☐ Requested strength/dose is not commercially available ☐ Other:						
this review?						

<u>Please note</u>: This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Otrexup® Prior Authorization Request Form

	OO NOT COPY FOR FUT	URE USE. FORMS ARE U	PDATED FREQUENTLY	AND MAY BE	BARCODED	
Memb	er Information	(required)	Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#: Specialty:			
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:	<u>l</u>	L	City:	State:		Zip:
	N	ledication Info	rmation (required)		
Medication Name:	•		Strength:	,	Dosage F	orm:
☐ Check if requesting b	orand		Directions for Use:			
☐ Check if request is fo	r continuation of ther	ару	_			
		Clinical Inform	nation (required)			
For active polyartics following: Is the patient intolera Has the patient tried at 180 days? Yes	r juvenile idiopathic assumatoid arthritis (RAnt, disabling psoriasis ular juvenile idiopatent of or has had an in and failed one month)	r severe, active rhe ofirst-line therapy? [form of methotrexat	umatoid ar	thritis (RA Io	
Has the patient had in	nadequate response and failed one month	to other forms of thera of a standard dosage	apy? ☐ Yes ☐ No	e (e.g., oral	, injectable) within the last
Are there any other com this review?	ments, diagnoses, symp	otoms, medications tried o	or failed, and/or any oth	er information	the physicia	an feels is important to
Please note: This	equest may be denied unl	ess all required information	is received			

For urgent or expedited requests please call 1-855-401-4262.



Praluent® & Repatha® Prior Authorization Request Form

Member Information (required)			Provider Information (required)			
Member Name:			Provider Nam	ne:		
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:	:	<u> </u>	
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street	Office Street Address:		
Phone:		<u>I</u>	City:	State:	Zip:	
		Medication	n Information	(required)		
Medication Name:			Strength:	(roquirou)	Dosage Form:	
☐ Check if requesting	g brand		Directions for	Use:		
☐ Check if request is	for continuatio	n of therapy				
		Clinical I	nformation (re	quired)		
☐ Homozygous far ☐ Hyperlipidemia i ☐ Other diagnosis: Clinical information Is the patient's base Has the patient been 80 mg, rosuvastation Is the patient a non- rhabdomyolysis or a normal [ULN])? ☐ Is the requested me	emilial hyperchemilial hyperchemilial hyperchemical high risk primary in a high risk primary in the LDL-C level receiving high tab 20 mg, ronguscle symptomuscle	suvastatin tab 40 mg)? high dose statin therap	al to 70 mg/dL? Yes No y (e.g., labeled cont nt with creatine kinas	Yes □ No (i.e., atorvastatin raindication to all see elevations great	tab 40 mg, atorvastatin tab statins, patient has experienced ater than 10 times upper limit of	
Is there documenta baseline? Yes	tion of positive	· 	erapy with LDL level		dl or decreased 30% from on the physician feels is important to	

Please note: This request may be denied unless all required information is received.



Proton Pump Inhibitor Prior Authorization Request Form

	DO NOT COPY FOR FUT	URE USE. FORMS ARE U	IPDATED FREQU	ENTLY AND MAY BE	BARCODED
Memb	er Information	(required)	P	rovider Info	rmation (required)
Member Name:			Provider Name	e:	
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone:		
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street A	Address:	
Phone:		<u> </u>	City:	State:	Zip:
		Medication Inf	ormation (required)	
Medication Name:		modioation iii		required	Danaga Form:
			Strength:		Dosage Form:
☐ Check if requesting	brand for continuation of the	rany	Directions for	Use:	
a officer if request is	or continuation of the	Clinical Infor	mation (rec	uirod\	
Select the diagnosis	holowy		mation (rec	quirea)	
☐ Barrett's esophagit		esophagitis	□ Zollinger-E	Ilison Syndrome	
Other diagnosis:	is 🗖 Elosive	esopriagitis	_	D-10 Code(s):	
release suspension per the following:	pack, Protonix packet,		et (omeprazole	/sodium bicarbona	let [ODT]), Prilosec delayed ite oral packet) requests, answer
		•			a) Provincely and neels
	cillin-clarithromycin o				e), Prevpack oral pack cole-sodium bicarbonate
	trial and failure (after a socie, or rabeprazole?		the past year wit	th at least one of the	following generics: Lansoprazole,
Has the patient experi- following: Lansoprazol	enced an adverse react le, omeprazole, pantopr	ion (must be documente azole, and rabeprazole?	ed on a MedWate You Yes D No	ch form), allergy or o	contraindication to <u>ALL</u> of the
Quantity limit reques What is the quantity re	sts: equested per DAY?				
	or exceeding the plan				
Titration or loadingPatient is on a dose	dose purposes	e.g., one tablet in the mo	orning and two ta	ablets at night, one t	to two tablets at bedtime)
Are there any other conthis review?	nments, diagnoses, sym _l	otoms, medications tried	or failed, and/or a	any other information	n the physician feels is important to
					
Please note: This	request may be denied un	less all required information	n is received.		



Duexis® & Vimovo® Prior Authorization Request Form (Page 1 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

	er Informa	tion (required)			rmation (required)
Member Name:			Provider Nan	ne:	
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone	:	
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street Address:		
Phone:		L	City:	State:	Zip:
		Medication In	formation	(required)	
Medication Name:			Strength:	(.oquou)	Dosage Form:
☐ Check if requesting	brand		Directions for	r Use:	
☐ Check if request is	for continuation	of therapy			
		Clinical Info	rmation (re	equired)	
Select the diagnos	sis below:				
☐ Ankylosing spon		only]			
Osteoarthritis					
□ Rheumatoid arth	nritis				
☐ Other diagnosis:	:		ICD-10) Code(s):	
Clinical information	n:				
Does the patient ha	ive a history of p	peptic ulcer disease/gastr	ointestinal (GI)	bleed? ☐ Yes ☐	l No
Does the patient hat corticosteroids)?		al risk factor for gastroint	estinal adverse	events (e.g., use	of anticoagulants, chronic
Does the patient ha	ive a history of a	asthma or urticaria after ta	aking aspirin or	other NSAIDs?	⊒Yes □ No
For Duexis reques	ts, please also	answer the following:			
		f a preferred generic H2- vithin the last 180 days?		er (e.g., famotidine	, cimetidine, ranitidine,
For Vimovo reque	sts, please also	o answer the following:			
		f a preferred generic prot D within the last 180 day			ole, lansoprazole,
Quantity limit requ					
What is the quantity					
		the plan limitations?			
tablets at bedtim	lose-alternating ne)	schedule (e.g., one table	t in the morning	g and two tablets a	at night, one to two
☐ Requested strer☐ Other:		commercially available			



Duexis® & Vimovo® Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

this review?	ther comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to
Please note:	This request may be denied unless all required information is received. For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.



Qualaquin® (quinine) Prior Authorization Request Form

	DO NOT COPY F	OR FUTURE USE. FORMS A	RE UPDATED FREQU	JENTLY AND MAY BI	E BARCODED
Memb	oer Informa	ation (required)			rmation (required)
Member Name:			Provider Nam	e:	
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone:		
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street	Address:	
Phone:			City:	State:	Zip:
		Medication	Information	(required)	
Medication Name:			Strength:	(,	Dosage Form:
☐ Check if requesting	g brand		Directions for	Use:	
☐ Check if request is	for continuation	of therapy			
		Clinical In	formation (red	quired)	
Select the diagn	osis below:				
☐ Malaria					
Other diagnos	is:		ICD-10 Co	ode(s):	
Quantity limit re	•	DAYO			
•	•	per DAY?			
■ Titration or loa		ding the plan limitation	ons?		
			e tablet in the mo	orning and two ta	ablets at night, one to two
tablets at bedt		9 (- 9 , -		9	3 4, 2 2 2 2
		not commercially avai			
U Other:					
Are there any other co this review?	mments, diagnose	es, symptoms, medications	tried or failed, and/or	any other informatio	n the physician feels is important to
		enied unless all required inform			



Rayos® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Р	rovider Info	rmation (required)		
Member Name	e :		Provider Name	e:			
Insurance ID#:			NPI#:	Specialty:			
Date of Birth:			Office Phone:				
Street Address	3:		Office Fax:				
City:	State:	Zip:	Office Street A	Address:			
Phone:	I	I	City:	State:	Zip:		
		Medication I	nformation	required)			
Medication Na	me:		Strength:	Strength: Dosage Form:			
☐ Check if req	uesting brand		Directions for	Directions for Use:			
☐ Check if req	uest is for continuatio	n of therapy					
		Clinical In	formation (red	quired)			
Has the pati	ient had a trial and	failure of generic predn	isone tablets in t	the past 60 days	? 🗆 Yes 🗅 No		
Are there any or this review?	ther comments, diagnos	es, symptoms, medications t	ried or failed, and/or	any other information	n the physician feels is important to		
Please note:	. ,	enied unless all required inform					



Relistor® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Pr	Provider Information (required)			
Member Name:			Provider Name) :			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:		1		
Street Address:			Office Fax:	Office Fax:			
City:	State:	Zip:	Office Street A	ddress:			
Phone:	I	L	City:	State:	Zip:		
		Medication	Information (red	quired)			
Medication Name:			Strength:		Dosage Form:		
☐ Check if request	•		Directions for U	Jse:	L		
☐ Check if request	t is for continuation o	f therapy					
		Clinical In	nformation (requir	red)			
•	ed constipation in adsis:	lult patients with adva	nced illness ICD-10) Code(s):			
•			e other laxative (e.g.,	stimulant, osmoti	ic, bulk forming, etc.) in the		
Are there any other this review?	er comments, diagnose	es, symptoms, medication	ns tried or failed, and/or a	nny other information	n the physician feels is important to		
Please note:	For urgent or expedited	enied unless all required int d requests please call 1-85 l for non-urgent requests ar).			

141



Soma® 250 (carisoprodol) Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Member Information (required)			Provider Information (required)				
Member Name:			Provider Name:				
Insurance ID#:			NPI#: Specialty:				
Date of Birth:			Office Phone:		II.		
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Address:				
Phone:			City:	State:		Zip:	
		Medication Inf	ormation (required	1)			
Medication Name:			Strength:	<u></u>	Dosage Fo	orm:	
☐ Check if requesting	brand		Directions for Use:				
☐ Check if request is f	or continuation of the	rapy					
		Clinical Infor	mation (required)				
Select the diagno	sis below:						
Acute painful m	nusculoskeletal con	dition					
Other diagnosis	S:		ICD-10 Code(s):				
Medication histor							
Has the patient ha	d a 6 month trial of	carisoprodol 350 m	g within the last 120	days? 🗖	Yes 🗆 N	lo	
Quantity limit req							
· •	ty requested per DA						
	on for exceeding the ding dose purposes	ne plan limitations	?				
			blet in the morning a	and two ta	blets at nic	aht, one to two	
tablets at bedting		rioddio (orgi, orio ta	order in the morning t		Dioto at III,	g, σσ το τπο	
□ Requested street	,	mmercially available	е				
Other:							
Are there any other con this review?	nments, diagnoses, sym _l	otoms, medications tried	or failed, and/or any othe	r information	the physicia	n feels is important to	
							

<u>Please note</u>: This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Tivorbex[™] Prior Authorization Request Form

Member Information (required)			P	Provider Information (required)				
Member Name	:		Provider Name	Provider Name:				
Insurance ID#:			NPI#: Specialty:					
Date of Birth:			Office Phone:					
Street Address:	:		Office Fax:					
City:	State:	Zip:	Office Street A	ddress:				
Phone:	I	I	City:	State:	Z	Zip:		
		Medication	Information (required)				
Medication Nar	me:		Strength:					
☐ Check if requ	uesting brand		Directions for U	Jse:				
☐ Check if requ	uest is for continuation	of therapy						
		Clinical Ir	nformation (req	uired)				
		ailure (a minimum of ory drugs (NSAIDs				iption strength		
Are there any ot this review?	ther comments, diagnose	es, symptoms, medications	s tried or failed, and/or a	any other information	n the physician f	feels is important to		
Please note:	, ,	enied unless all required info I requests please call 1-855-						



Conzip[®], Synapryn[®], Ultram[®] ER (tramadol ER biphasic capsule or tablet) Prior Authorization Request Form

D	O NOT COPY FOR FUTU	IRE USE. FORMS ARE UF	DATED FREQUENTLY A	ND MAY BE	BARCODED		
Memb	er Information	(required)	Provider Information (required)				
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Address:				
Phone:		1	City:	State:		Zip:	
		Medication Info	rmation (required)				
Medication Name:			Strength:		Dosage Fo	orm:	
☐ Check if requesting	brand		Directions for Use:				
☐ Check if request is	for continuation of the	rapy					
		Clinical Inform	nation (required)				
Clinical information:							
Is the patient currently	stable on tramadol ER	tablet or Ultram ER? □	l Yes □ No				
Is the patient currently	stable on Conzip, Syna	apryn (tramadol suspens	sion), tramadol ER bipha	asic capsule	or tablet?	⊒Yes □ No	
Has the patient failed	a 30 day trial of immedi	ate release tramadol in t	the last 120 days? 🛚 Y	es 🛭 No			
Has the patient had ar form? ☐ Yes ☐ No	n adverse reaction to ge	eneric immediate-release	e tramadol and the pres	criber has do	ocumented it	t on a MedWatch	
Has the patient had a patient's chart notes/m	drug allergy or contrain nedical records? ☐ Yes	dication to generic imme D No	ediate-release tramadol	and the pres	scriber has d	locumented it in the	
Does the patient have	a diagnosis of cancer in	n the past 365 days? 🛚	Yes □ No				
Does the patient have	a diagnosis of a termin	al illness? 🛭 Yes 🗖 No	o				
Does the patient have If yes , please list the o	· · · · · · · · · · · · · · · · · · ·	rith significant pain (e.g.	sickle cell anemia, etc)	? 🗆 Yes 🗆	l No		
Does the patient have If yes , please list the o		th significant pain? 🗖 Y	es 🗆 No				
Have efforts been mad If yes , please provide		the lowest effective do	se? 🛘 Yes 🗀 No				
Reauthorization:							
If this is a reauthoriz	ation request, answer	the following:					
•		vative, effective treatme					
If yes , please provide	documentation:						
Ara thara any other com	manta diampaga aumot	ome medications tried a	r failed and/or any other	information	the physicial	n faala ia immawtant ta	

Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?

<u>Please note:</u> This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Triptans Prior Authorization Request Form

Memb	er Informa	tion (required)			rmation (required)		
Member Name:			Provider Name	:			
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:	Office Fax:			
City:	State:	Zip:	Office Street Ad	Office Street Address:			
Phone:	l .	-	City:	State:	Zip:		
		Medication	n Information (r	equired)			
Medication Name:			Strength:	,	Dosage Form:		
☐ Check if requesting	brand		Directions for U	lse:	<u> </u>		
☐ Check if request is	for continuation	of therapy					
		Clinical I	nformation (requ	ıired)			
Select the diagno	osis below:						
☐ Migraine with o	or without aura	1					
Other diagnosi	s:		ICD-	-10 Code(s):			
Medication histo	ry:						
Has the patient ha	ad a trial and f	ailure of a generic	triptan within the las	t 6 months?	I Yes □ No		
Clinical informat	ion:						
Does the patient h	nave a diagnos	sis which confirms	a difficulty in swallo	wing? \(\sigma\) Yes	□ No		
Quantity limit red							
What is the quanti	• •						
		ling the plan limit	ations?				
☐ Titration or load ☐ Patient is on a			one tablet in the mo	orning and two	tablets at night, one to two		
tablets at bedti		ing contodulo (c.g.,		ining and two	tablete at mgm, one to two		
•	ength/dose is i	not commercially a	vailable				
☐ Other:							
Are there any other com	ments, diagnoses	, symptoms, medication	s tried or failed, and/or a	ny other information	n the physician feels is important to		

This request may be denied unless all required information is received. Please note:

For urgent or expedited requests please call 1-855-401-4262.



Onzetra® Xsail® Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Mo	ember Inform	ation (required)	Р	rovider Info	rmation (required)	
Member Name	:		Provider Nam	ne:		
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone	Phone:		
Street Address:	:		Office Fax:			
City:	State:	Zip:	Office Street	Address:		
Phone:			City:	State:	Zip:	
		Medication In	formation	(required)		
Medication Name:			Strength: Dosage Form:		Dosage Form:	
☐ Check if requ	uesting brand		Directions for Use:			
☐ Check if requ	uest is for continuatio	on of therapy				
		Clinical Info	rmation (re	equired)		
Has the patie	ent had a trial and	failure to at least six other	er triptans in t	the past 36 mont	ths? 🛘 Yes 🗘 No	
Are there any oth this review?	er comments, diagnos	es, symptoms, medications tried	l or failed, and/or	any other informatio	on the physician feels is important to	
Please note:	This request may be d	enied unless all required information	on is received.			

For urgent or expedited requests please call 1-855-401-4262.



Nurtec ODTTM, QuliptaTM, Reyvow[®], UbrelvyTM Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

	er Information	(required)	Provide			required)
Member Name:			Provider Name:			
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:			
Street Address:			Office Fax: Office Street Address:			
City:	State:	Zip:	Office Street Address:			
Phone:		1	City:	State:		Zip:
	N	ledication Info	mation (required)			
Medication Name:			Strength:		Dosage Fo	orm:
☐ Check if requesting			Directions for Use:			
☐ Check if request is	for continuation of the					
		Clinical Inform	nation (required)			
Preventive treatment	of migraine with or winent of episodic migra		ICD-10 Coo	de(s):		
Clinical information						
Has the patient had	a trial and failure of a	triptan in the last 120	days? 🛚 Yes 🗖 No			
•	·	nse, intolerance to, or	contraindication to tri	ptans? 🗖	Yes 🛭 No	
-	ve cardiovascular dis	ease? Yes No				
What is the reason ☐ Titration or loadin ☐ Patient is on a do ☐ bedtime) ☐ Requested stren	requested per DAY? for exceeding the page dose purposes ose-alternating schedulers and the page of the	olan limitations?	-	tablets at r	night, one t	o two tablets at
		toms, medications tried o		information t	the physiciar	n feels is important

<u>Please note:</u> This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Uloric Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Memb	per Informatio	n (required)	Provi	der Info	rmatior	(required)		
Member Name:			Provider Name:					
Insurance ID#:			NPI#:		Specialty:			
Date of Birth:			Office Phone:		l.			
Street Address:			Office Fax:					
City:	State:	Zip:	Office Street Address:					
Phone:	1	I	City:	State:		Zip:		
		Medication Inf	ormation (require	ed)				
Medication Name:			Strength:	,	Dosage F	orm:		
☐ Check if requesting		Directions for Use:						
☐ Check if request is	for continuation of the	erapy						
		Clinical Infor	mation (required)					
Select the diagno	osis below:							
□ Chronic gout								
Other diagnosi	s:		_ ICD-10 Code(s)	:				
Clinical informat	ion:							
Has the patient re	ceived an adequat	e trial of at least 1 m	onth of allopurinol?	? 🗆 Yes 🗆	l No			
Does the patient h	nave renal or hepat	ic dysfunction? 🗖 Y	es 🛭 No					
Are there any other couthis review?	mments, diagnoses, syn	nptoms, medications tried	or failed, and/or any oth	ner information	the physici	an feels is important to		
	. ,	nless all required information						



ViberziTM Prior Authorization Request Form

Memb	er Informat	ion (required)			mation (require	ed)
Member Name:			Provider Name:		(,	
Insurance ID#:			NPI#:		Specialty:	
Date of Birth:			Office Phone:		I.	
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Addre	ess:		
Phone:			City:	State:	Zip:	
		Medication Inf	ormation (requi	ired)		
Medication Name:			Strength:		Dosage Form:	
☐ Check if requesting			Directions for Use:			
☐ Check if request is	for continuation o f					
		Clinical Infor	mation (required	d)		
Select the diagno		diarrhea (IBS-D)				
	•		_ ICD-10 Code(s	s):		
Are there any other cor this review?	nments, diagnoses,	symptoms, medications tried	or failed, and/or any o	ther information	the physician feels i	s important to
Please note: This	request may be denie	ed unless all required informatio	n is received.			

For urgent or expedited requests please call 1-855-401-4262. This form may be used for non-urgent requests and faxed to 1-844-403-1029.



Xenazine® Prior Authorization Request Form

DO NOT COPY FOR FUTURE USE, FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

	lember Inform	ation (required)			rmation (required)
Member Name	: :		Provider Nam	ie:	
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone:		
Street Address	3:		Office Fax:		
City:	State:	Zip:	Office Street	Address:	
Phone:	l .		City:	State:	Zip:
		Medication	Information	(required)	
Medication Nar	me:		Strength:		Dosage Form:
☐ Check if req☐ Check if req☐	uesting brand uest is for continuatio r	of therapy	Directions for	Use:	1
			formation (re	quired)	
	ent have a confirmed	diagnosis of chorea ass		•	
-	· · · · · · · · · · · · · · · · · · ·	-	-		n the physician feels is important to
Please note:		enied unless all required inform			

For urgent or expedited requests please call 1-855-401-4262.



Xepi[™] Prior Authorization Request Form DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Men	nber Inform	nation (required)	F	Provider Info	ermation (required)
Member Name:			Provider Nam	ne:	
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone	:	
Street Address:		Office Fax:			
City:	State:	Zip:	Office Street	Address:	
Phone:		I	City:	State:	Zip:
		Medication	Information	(required)	
Medication Name:			Strength:		Dosage Form:
☐ Check if request	ting brand		Directions for	Use:	
□ Check if request	t is for continuatio	n of therapy			
		Clinical In	nformation (re	equired)	
	to Staphylococcu	us aureus or Streptococo		CD-10 Code(s):	
Medication histo				()	
Has the patient h	ad a 10 day trial	and failure of mupirocin	ointment/cream wi	thin the past 6 mo	nths?
Are there any other this review?	comments, diagnos	ses, symptoms, medications	s tried or failed, and/or	any other informatio	n the physician feels is important to
Please note:	This request may be o	denied unless all required info			

For urgent or expedited requests please call 1-855-401-4262.



Xifaxan® Prior Authorization Request Form

Memb		ation (required)			rmation (required)
Member Name:			Provider Nam	e:	
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone:		1
Street Address:			Office Fax:		
City:	State:	Zip:	Office Street A	Address:	
Phone:			City:	State:	Zip:
		Medication I	nformation	(required)	
Medication Name:			Strength:		Dosage Form:
☐ Check if requesting			Directions for	Use:	
☐ Check if request is	for continuatior	of therapy			
		Clinical Info	ormation (red	quired)	
Select the diagno	osis below:				
Hepatic encep	halopathy (HE	≣)			
□ Irritable bowel	syndrome wit	h diarrhea (IBS-D)			
Travelers' diar	rhea				
Other diagnosi	s:		ICD-10 Co	ode(s):	
Are there any other couthis review?	mments, diagnos	es, symptoms, medications tri	ed or failed, and/or	any other information	n the physician feels is important to
Please note: This	request may be d	enied unless all required informa	ition is received.		

For urgent or expedited requests please call 1-855-401-4262.



Ambien CR[®], Edluar[™], Intermezzo[®] (zolpidem sublingual tablet [SL]), Zolpimist[™] **Prior Authorization Request Form**

		or future use. Forms ar ation (required)			ermation (required)
Member Name:			Provider Nam		
Insurance ID#:			NPI#:		Specialty:
Date of Birth:			Office Phone:	:	
Street Address:	ss: Office Fax:				
City:	State:	Zip:	Office Street	Address:	
Phone:			City:	State:	Zip:
		Medication I	nformation	(required)	
Medication Name:		Medication	Strength:	(required)	Dosage Form:
☐ Check if requestin	g brand		Directions for	Use:	ŭ
☐ Check if request is	*	n of therapy			
		Clinical Inf	ormation (re	guired)	
Select the diagn	osis below:		,	·	
☐ Insomnia					
☐ Other diagnos	sis:		ICD-10 C	Code(s):	
reaction (prescrib	ad a trial (at le per must have	east a 14 day trial in the documented it on a Me r brand Ambien tablets	edWatch form),	or contraindica	e response, adverse tion to generic immediate
Quantity limit re What is the quan		per DAY?	_		
		ding the plan limitation	ons?		
tablets at bedi	a dose-alterna time) rength/dose is		lable	norning and two	tablets at night, one to two
				any other information	on the physician feels is important to

Please note:

This request may be denied unless all required information is received.

For urgent or expedited requests please call 1-855-401-4262.



Belsomra[®], Dayvigo[®], Quviviq[™] Prior Authorization Request Form

Memb	er Informa	ntion (required)	Pro	ovider Infor	mation (required)		
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:	Office Fax:			
City:	State:	Zip:	Office Street Ad	Office Street Address:			
Phone:			City:	State:	Zip:		
		Medication	Information (re	equired)			
Medication Name:			Strength:		Dosage Form:		
☐ Check if requesting	brand		Directions for U	se:			
☐ Check if request is		of therapy					
		Clinical I	nformation (requ	ired)			
Select the diagno	osis below:						
☐ Insomnia							
☐ Other diagnosi	s:		ICD-10 Co	de(s):			
Medication histo	ry:						
			the last 180 days) a				
				r contraindicati	on to generic immediate		
·		brand Ambien table	ets? LI Yes LI No				
Quantity limit red		nor DAV2					
What is the quanti	•	ling the plan limita					
☐ Titration or load			1110115 ?				
			one tablet in the mo	orning and two t	ablets at night, one to two		
tablets at bedti	me)			•			
		not commercially av					
Uther:							
Are there any other com	ments, diagnoses	, symptoms, medications	s tried or failed, and/or an	y other information	the physician feels is important to		
his review?							
Please note: This re	equest may be den	ied unless all required info	rmation is received				

For urgent or expedited requests please call 1-855-401-4262.



Therapeutic Class Overview Attention-Deficit/Hyperactivity Disorder (ADHD) Agents

INTRODUCTION

- Attention-deficit/hyperactivity disorder (ADHD) is the most common neurodevelopmental disorder among children, with an estimated prevalence of up to 10% in school-age children in the United States (US). It is more common in boys than girls and frequently persists into adulthood (*Centers for Disease Control and Prevention [CDC] 2021, Feldman et al* 2014). Epidemiologic studies of adult ADHD have estimated the current prevalence to be 4.4% in the US. (*Bukstein* 2022).
 - o In children, this chronic disorder is characterized by symptoms of hyperactivity, impulsivity, and/or inattention. These symptoms affect cognitive, academic, behavioral, emotional, and social functioning (*Krull 2022a*). Common comorbid psychiatric disorders include oppositional defiant disorder, conduct disorder, depression, anxiety disorder, and learning disabilities (*Krull 2022b*). Approximately 20% of children with ADHD develop chronic tic disorders and approximately 50% of children with chronic tics or Tourette syndrome have comorbid ADHD (*Krull 2022c*).
 - ADHD in adults is characterized by symptoms of inattention, impulsivity, and restlessness. Impairment in executive
 function and emotional dysregulation frequently occur. Common comorbid psychiatric disorders include mood and
 anxiety disorders, substance use disorder, and intermittent explosive disorder (*Bukstein 2022*).
- For children < 17 years of age, the Diagnostic and Statistical Manual of Mental Disorders Fifth Edition (DSM-5) diagnosis of ADHD requires ≥ 6 symptoms of hyperactivity and impulsivity or ≥ 6 symptoms of inattention. For adolescents ≥ 17 years of age and adults, ≥ 5 symptoms of hyperactivity and impulsivity or ≥ 5 symptoms of inattention are required.
 - The symptoms of hyperactivity/impulsivity or inattention must occur often; be present in more than 1 setting; persist
 for at least 6 months; be present before the age of 12 years; impair function in academic, social, or occupational
 activities; and be excessive for the developmental level of the child.
 - o Other physical, situational, or mental health conditions that could account for the symptoms must be excluded.
- Treatment of ADHD may involve behavioral/psychologic interventions, medication, and/or educational interventions, alone or in combination (*Krull 2022d*).
 - o For preschool children (age 4 through 5 years), behavioral therapy is considered the first-line treatment; when medication is necessary, methylphenidate is generally recommended.
 - For children and adolescents with moderate to severe ADHD, medication and behavioral therapy are recommended. In general, stimulants are the first-line agents; however, nonstimulant medications may be more appropriate for certain children.
 - Some patients do not respond to or may not tolerate the initial stimulant treatment. At least one-half of children who do not respond to one type of stimulant will respond to the other. If there is still no improvement, consideration should be given to switching to or adding a nonstimulant ADHD medication (*Krull 2022e*).
- Multiple agents are currently approved by the Food and Drug Administration (FDA) for the treatment of ADHD. They include central nervous system (CNS) stimulants (amphetamine- and methylphenidate-based formulations), as well as nonstimulants: 2 selective norepinephrine reuptake inhibitors (SNRIs), atomoxetine and viloxazine extended-release (ER); and 2 alpha₂-adrenergic agonists, clonidine ER and quanfacine ER.
 - o Due to the potential for abuse, the stimulant agents are classified as Schedule II controlled substances.
 - Several stimulants are also approved for the treatment of narcolepsy and exogenous obesity; the use of stimulants for the treatment of obesity will not be covered in this review. Lisdexamfetamine dimesylate is the only FDA-approved drug for the treatment of binge eating disorder (BED).
- Medispan Classes: ADHD Agents Amphetamines, Dexmethylphenidate, Methylphenidate, Selective Alpha-Adrenergic Agonists, Selective Norepinephrine Reuptake Inhibitor



Table 1. Medications Included Within Class Review

Drug	Generic Availability
Stimulants	
Evekeo (amphetamine sulfate)	✓
Evekeo ODT (amphetamine sulfate)	-
Azstarys (serdexmethylphenidate/dexmethylphenidate)	-
Adderall (mixed amphetamine salts)	∨
Focalin (dexmethylphenidate hydrochloride [HCI])	∨
ProCentra (dextroamphetamine sulfate)	∨
Zenzedi (dextroamphetamine sulfate)	∨
Xelstrym (dextroamphetamine transdermal system)	-
Desoxyn (methamphetamine HCl)	✓
methylphenidate HCl chewable tablets	✓
Methylin Oral Solution (methylphenidate HCI)	✓
Ritalin (methylphenidate HCI)	✓
Dexedrine Spansule (dextroamphetamine sulfate	,
sustained-release)	•
Adzenys XR-ODT (amphetamine ER)	-
Dyanavel XR (amphetamine ER)	-
Adderall XR (mixed amphetamine salts ER)	✓
Mydayis (mixed amphetamine salts ER)	<u>-</u>
Focalin XR (dexmethylphenidate HCl ER)	✓
Vyvanse (lisdexamfetamine dimesylate)	<u>-</u>
Adhansia XR (methylphenidate HCl ER)*	<u>-</u>
Aptensio XR (methylphenidate HCI ER)	✓
Concerta (methylphenidate HCI ER)	✓
Cotempla XR-ODT (methylphenidate ER)	-
Jornay PM (methylphenidate HCl ER)	-
methylphenidate HCl ER (CD)	✓
methylphenidate HCl ER	✓
QuilliChew ER (methylphenidate HCl ER)	<u>-</u>
Quillivant XR (methylphenidate HCl ER)	<u>-</u>
Relexxii (methylphenidate HCl ER) (72 mg)	✓
Ritalin LA (methylphenidate HCl ER)	✓
Daytrana (methylphenidate transdermal system)	<u> </u>
Nonstimulants	
Strattera (atomoxetine HCI)	✓
Kapvay (clonidine HCl ER)	✓
Intuniv (guanfacine HCl ER)	✓
Qelbree (viloxazine ER)	<u>-</u>
*Adhansia XR was discontinued by the manufacturer in July 2022	

*Adhansia XR was discontinued by the manufacturer in July 2022.

(Drugs@FDA 2022, Orange Book: Approved Drug Products with Therapeutic Equivalence Evaluations 2022, Clinical Pharmacology 2022)



INDICATIONS

Table 2. Food and Drug Administration Approved Indications

Indication	ADHD*	ADHD, as an integral part of a total treatment program which typically includes other remedial measures (psychological, educational, and social) for a stabilizing effect in pediatric patients with a behavioral syndrome characterized by the following group of developmentally inappropriate symptoms: moderate to severe distractibility, short attention span, hyperactivity, emotional lability, and impulsivity. The diagnosis of this syndrome should not be made with finality when these symptoms are only of comparatively recent origin. Nonlocalizing (soft) neurological signs, learning disability, and abnormal electroencephalogram (EEG) may or may not be present, and a diagnosis of CNS dysfunction may or may not be warranted.*	Treatment of ADHD as monotherapy and as adjunctive therapy to stimulant medications	Narcolepsy**	Exogenous obesity, as a short term (a few weeks) adjunct in a regimen of weight reduction based on caloric restriction for patients refractory to alternative therapy (eg, repeated diets, group programs, and other drugs).†	Moderate to severe BED in adults
Evekeo (amphetamine sulfate)		√ ·		√	√	
Evekeo ODT (amphetamine sulfate)	✓					
Adzenys XR-ODT, Dyanavel XR (amphetamine)	✓					
Adderall (mixed amphetamine salts)	✓			√		
Adderall XR, Mydayis (mixed amphetamine salts ER)	√					
Strattera (atomoxetine HCl)	✓					
Kapvay (clonidine HCl ER)			✓			
Focalin (dexmethylphenidate IR); Focalin XR (dexmethylphenidate ER)	√					
ProCentra, Zenzedi (dextroamphetamine sulfate IR); Dexedrine Spansule (dextroamphetamine sulfate SR)		✓		√		
Intuniv (guanfacine HCI ER)			✓			
Vyvanse (lisdexamfetamine dimesylate)	\checkmark	i		1		✓

Data as of October 11, 2022 HJ-U/KS-U/AVD

Page 157
This information is considered confidential and proprietary to OptumRx. It is intended for internal use only and should be disseminated only to authorized recipients. The contents of the therapeutic class overviews on this website ("Content") are for informational purposes only. The Content is not intended to be a substitute for professional medical advice, diagnosis, or treatment. Patients should always seek the advice of a physician or other qualified health provider with any questions regarding a medical condition. Clinicians should refer to the full prescribing information and published resources when making medical decisions.



Desoxyn (methamphetamine HCI)		✓		
Ritalin (methylphenidate HCl IR); methylphenidate HCl chewable tablets		✓	✓	
Methylin Oral Solution; methylphenidate ER tablets	✓		✓	
Adhansia XR, Aptensio XR, Concerta, Cotempla XR-ODT, Daytrana, Jornay PM, QuilliChew ER, Quillivant XR, Relexxii, Ritalin LA (methylphenidate ER)	√			
Azstarys (serdexmethylphenidate)	✓			
Qelbree (viloxazine ER)	√			
Xelstrym (dextroamphetamine transdermal)	✓			

(Prescribing Information: Adderall 2022, Adderall XR 2022, Adhansia XR 2021, Adzenys XR-ODT 2022, Aptensio XR 2021, Azstarys 2021, Concerta 2022, Cotempla XR-ODT 2021, Daytrana 2021, Desoxyn 2019, Dexedrine Spansule 2022, Dyanavel XR 2022, Evekeo 2022, Evekeo ODT 2021, Focalin 2021, Focalin XR 2021, Intuniv 2020, Jornay PM 2021, Kapvay 2020, Mydayis 2022, Methylin Oral Solution 2021, methylphenidate chewable tablets 2021, methylphenidate ER 2021, methylphenidate ER 2021, methylphenidate ER 2021, Ritalin 2021, Ritalin LA 2021, Strattera 2022, Vyvanse 2022, Xelstrym 2022, Zenzedi 2022)

*Adderall, Evekeo, ProCentra, and Zenzedi are approved for use in children 3 years of age and older. Evekeo ODT is approved for use in patients 3 to 17 years of age. Daytrana, Desoxyn, Dexedrine Spansule, Intuniv, and Kapvay are approved for use in children 6 years of age and older. Adderall XR, Adhansia XR, Adzenys XR-ODT, Aptensio XR, Azstarys, Dyanavel XR, Focalin, Focalin XR, Jornay PM, methylphenidate ER (CD), methylphenidate ER, Methylin Oral Solution, methylphenidate chewable tablets, Qelbree, QuilliChew ER, Quillivant XR, Ritalin, Strattera, Vyvanse, and Xelstrym are approved for use in patients 6 years of age and older. Cotempla XR-ODT is approved for use in pediatric patients 6 to 17 years of age. Ritalin LA is approved for use in pediatric patients 6 to 12 years of age. Concerta and Relexxii are approved for use in children 6 years of age and older, adolescents, and adults up to 65 years of age. Mydayis is approved for use in patients 13 years of age and older.

**These drugs are approved for use in patients 6 years of age and older.

†These drugs are not recommended for use in children under 12 years of age for treatment of exogenous obesity. The limited usefulness of these products should be weighed against possible risks inherent in use of the drugs.

- Limitation of use:
 - Aptensio XR: Pediatric patients younger than 6 years of age experienced higher plasma exposure than patients 6 years and older at the same dose and high rates of adverse events (AEs), most notably weight loss.
 - Lisdexamfetamine: Pediatric patients younger than 6 years of age experienced more long-term weight loss than
 patients 6 years and older. Lisdexamfetamine is not indicated or recommended for weight loss. Use of other
 sympathomimetic drugs for weight loss has been associated with serious cardiovascular (CV) AEs. The safety and
 effectiveness of this drug for the treatment of obesity have not been established.
 - Mydayis: Pediatric patients 12 years and younger experienced higher plasma exposure than patients 13 years and older at the same dose and experienced higher rates of AEs, mainly insomnia and decreased appetite.
 - Xelstrym: Pediatric patients younger than 6 years of age experienced more long-term weight loss than patients 6
 years and older.
- Information on indications, mechanism of action, pharmacokinetics, dosing, and safety has been obtained from the prescribing information for the individual products, except where noted otherwise.

CLINICAL EFFICACY SUMMARY

 Randomized trials, systematic reviews, and meta-analyses have found stimulants, SNRIs (atomoxetine, viloxazine ER), and alpha₂-adrenergic agonists (clonidine ER, guanfacine ER) to be more efficacious than placebo in reducing the core symptoms of ADHD in children and adolescents.



- Evekeo (amphetamine sulfate) was approved based on a randomized, double-blind (DB), multicenter (MC), placebo-controlled (PC) laboratory classroom study that was conducted in 107 children between the ages of 6 and 12 years (*Childress et al 2015*). The study found Evekeo to be associated with significant improvements in the average Swanson, Kotkin, Agler, M-Flynn, and Pelham (SKAMP) combined score compared to placebo (least squares [LS] mean difference -7.9; 95% confidence interval [CI], -10.1 to -5.6; p < 0.0001).
 - Evekeo ODT, an orally disintegrating amphetamine tablet, was approved under the 505(b)(2) regulatory pathway. The safety and effectiveness of Evekeo ODT for the treatment of ADHD was established based on an adequate and well-controlled study of Evekeo (*Childress et al 2015*).
- Cotempla XR-ODT, a new methylphenidate ER orally disintegrating tablet formulation, was approved based on a randomized, DB, MC, PC laboratory classroom study (*Childress et al 2017*) (N = 87) which found that the average SKAMP-combined score was significantly better for Cotempla XR-ODT than for placebo (LS mean 14.3 [95% CI, 12.2 to 16.4] vs 25.3 [9% CI, 23.0 to 27.6], respectively; p < 0.0001).
- Adhansia XR (methylphenidate ER capsule) was approved via the 505(b)(2) regulatory pathway, and its efficacy was supported by 4 clinical studies in patients with ADHD including 2 studies conducted in adults, 1 study in adolescents 12 to 17 years of age, and 1 study in pediatric patients 6 to 12 years of age (Adhansia XR FDA Clinical Review 2019):
 - One randomized, DB, MC, PC 4-week study conducted in 368 adult patients with ADHD evaluated the safety and efficacy of 4 doses of Adhansia XR (25, 45, 70, and 100 mg) compared to placebo. The primary endpoint, change in the ADHD-Rating Scale (ADHD-RS)-5 total score from baseline to Week 5, was significantly improved compared to placebo in the Adhansia XR 45 mg group (LS mean difference, -6.9; 95% CI, -11.5 to -2.2; p = 0.0013), 100 mg group (LS mean difference, -8.1; 95% CI, -12.9 to -3.2; p = 0.0002), and when combining all dosage groups compared to placebo (LS mean difference, -4.7; 95% CI, -7.7 to -1.6; p = 0.0026). No significant difference was seen in the 25 mg or 70 mg groups compared to placebo.
- o A second randomized, DB, crossover, PC study was conducted in 45 adults in an adult workplace environment (*Adhansia XR FDA Clinical Review 2019, Wigal et al 2020*). The study aimed to assess efficacy parameters for Adhansia XR vs placebo over 16 hours post-dose. Patients were titrated to an optimal dose of Adhansia XR (either 25, 35, 45, 55, 70, 85, or 100 mg) during an open-label (OL) treatment period between 2 and 7 weeks, then entered into a 1-week PC, DB treatment phase prior to the adult workplace environment session, followed by a 7-day washout period between crossover periods, then another 1-week treatment phase followed by another adult workplace environment session. The primary endpoint was the average Permanent Product Measure of Performance (PERMP) score for various time points up to 16 hours post-dose. When combining data from all time points, patients treated with Adhansia XR had significant improvements in the PERMP score compared to placebo (LS mean difference, 13.05; 95% CI, 3.88 to 22.23; p = 0.0064).
- o A 4-week randomized, DB, PC trial assessed efficacy of Adhansia XR in 354 adolescent patients 12 to 17 years of age (*Adhansia XR FDA Clinical Review 2019*). The study compared Adhansia XR 25, 45, 70, and 85 mg to placebo and found significant improvements in the ADHD-5-RS score from baseline to Week 5 in adolescents treated with Adhansia XR 45 mg (LS mean difference, -5.4; 95% CI, -9.2 to -1.6; p = 0.0052), 70 mg (LS mean difference, -5.2; 95% CI, -9.0 to -1.4; p = 0.0069), and when combining all dosage groups compared to placebo (LS mean difference, -4.3; 95% CI, -7.3 to -1.3; p = 0.0049). Adolescents treated with Adhansia XR 25 or 85 mg did not achieve significant improvements in the ADHD-5-RS score compared to placebo.
 - A fourth study, which included a 6-week OL dose optimization period (majority of patients received between 45 and 55 mg of Adhansia XR) followed by a 1- week DB, PC study, was conducted to assess the efficacy of Adhansia XR in 147 children 6 to 12 years of age in an analog classroom setting. The primary endpoint, average SKAMP-C score (taken at various time points up to 13 hours post-dose), was significantly improved in children treated with Adhansia XR compared to placebo (LS mean difference, -8.6; 95% CI, -10.6 to -6.6).
- o Jornay PM, an ER methylphenidate capsule formulation, was approved based on the results of 2 clinical studies conducted in patients 6 to 12 years of age with ADHD:
 - The first study was a 6-week OL dose-optimization study, followed by a 1-week DB, PC withdrawal phase where patients were randomized to continue treatment with Jornay PM or switch to placebo (*Childress et al 2020, Jornay PM Prescribing Information 2021*). The study, which was conducted in an analog classroom setting and included 117 children aged 6 to 12 years, found that Jornay PM was associated with a significant reduction in the SKAMP symptom score over a 12-hour period (LS mean difference, -5.9; 95% CI, -9.1 to -2.7).
 - A randomized, DB, MC, PC, parallel group, forced-dose titration trial was conducted over 3 weeks in 161 children 6 to 12 years of age with ADHD (*Pliszka et al 2017*). The study found that 40 to 80 mg/day of Jornay PM achieved significant improvements vs placebo in ADHD symptoms (LS mean ADHD rating scale-IV, 24.1 vs 31.2; p = 0.002)



- at 3 weeks. Significant improvements were also seen vs placebo in key secondary outcomes including at-home early morning and late afternoon/evening functional impairment at 3 weeks. The most commonly reported treatment-emergent AEs were insomnia and decreased appetite.
- o Mydayis, a mixed amphetamine salts product, was approved for the treatment of ADHD based on the results of 5 MC, DB, PC, randomized controlled trials (RCTs): 3 in adults and 2 in pediatric patients 13 to 17 years of age. The studies found that Mydayis demonstrated a statistically significant treatment effect compared with placebo on various ADHD outcomes measures (eg, ADHD-RS score, PERMP score) (*Mydayis Prescribing Information 2022, Weisler et al 2017, Wigal et al 2018a, Wigal et al 2018b, Wigal et al 2019*) (see results below in Table 3 below). An additional 6-week, randomized, PC, DB, forced dose titration trial in 411 adults with ADHD similarly found that Mydayis significantly improved ADHD-RS-IV scores compared to placebo (LS mean treatment difference for all Mydayis doses combined vs placebo, -10.6; 95% CI, -13.2 to -8.0; p < 0.0001) (*Frick et al 2020*).

Table 3. Summary of Primary Efficacy Results for Mydayis

Study Number	Primary Endpoint	Treatment Group	Mean Baseline Score (SD)	LS Mean Change	Placebo-subtracted Difference (95% CI)
(Age range)	Liidpoiit		33013 (32)	from Baseline	Billiotolioo (00% 01)
Adult Studies					
Study 1	ADHD-RS	Mydayis 12.5 mg/day [§]	39.8 (6.38)	-18.5	-8.1 (-11.7 to -4.4)
(18 to 55 years)		Mydayis 37.5 mg/day [§]	39.9 (7.07)	-23.8	-13.4 (-17.1 to -9.7)
,		Placebo	40.5 (6.52)	-10.4	
Study 2 (18 to 55	Average PERMP	Mydayis 50 mg/day [§]	239.2 (75.6)†	293.23*	18.38 (11.28 to 25.47)
years)		Placebo	249.6 (76.7) [†]	274.85*	
Study 3 (18 to 55	Average PERMP	Mydayis 25 mg/day [§]	217.5 (59.6)†	267.96*	19.29 (10.95 to 27.63)
years)		Placebo	226.9 (61.7) [†]	248.67*	
Pediatric Stud	ies				
Study 4 (13 to 17 years) [‡]	ADHD-RS-IV	Mydayis 12.5 to 25 mg/day [§]	36.7 (6.15)	-20.3	-8.7 (-12.6 to -4.8)
		Placebo	38.3 (6.67)	-11.6	
Study 5 (13 to 17	Average PERMP	Mydayis 25 mg/day [§]	214.5 (87.8)†	272.67*	41.26 (32.24 to 50.29)
years)		Placebo	228.7 (101) [†]	231.41*	

SD = standard deviation; LS = least squares; CI = confidence interval

- o Azstarys, a combination of serdexmethylphenidate and dexmethylphenidate, was approved based on results from a randomized, DB, PC analog classroom study (*Kollins et al 2021*). A total of 150 patients aged 6 to 12 years were enrolled. Following an OL, 3-week dose titration phase, patients were randomly assigned during a 1-week parallel treatment period to either the optimized dose Azstarys or placebo. After 1 week, evaluations were done using the SKAMP rating scale over 13 hours in a classroom setting. Mean change in SKAMP from baseline (primary outcome) was significantly greater with Azstarys compared with placebo (placebo-subtracted difference -5.41; 95% CI, -7.10 to -3.71; p < 0.001). The efficacy of Azstarys in adults and pediatric patients 13 to 17 years of age was established by pharmacokinetic bridging between Azstarys and Focalin XR (dexmethylphenidate ER) capsules.
- Qelbree (viloxazine ER), an SNRI, was shown to be superior to placebo in 3 DB, MC, randomized, PC trials in pediatric patients with ADHD.
 - Trial 1 enrolled 313 patients aged 6 to 11 years who were randomized to treatment with viloxazine ER 200 or 400 mg or placebo once daily for 8 weeks (*Nasser 2021b*). Improvements in ADHD-RS-5 total scores were reported, with LS mean changes from baseline of -17.6, -17.5 and -11.7 for viloxazine ER 200 mg, 400 mg, and placebo, respectively (p < 0.05 for both comparisons to placebo).</p>

[†]Pre-dose PERMP total score

^{*}LS mean for PERMP is post-dose average score over all sessions of the treatment day, rather than change from baseline

[‡]Results are for a subgroup of study 4 and not the total population

[§]Doses statistically significant for placebo



- Trial 2 enrolled 477 patients aged 6 to 11 years who were randomized to either viloxazine ER 100 mg or 200 mg or placebo once daily for 6 weeks (*Nasser 2020*). LS mean changes from baseline in ADHD-RS-5 total scores were -16.6, -17.7, and -10.9 for viloxazine ER 100 mg, 200 mg, and placebo, respectively (p < 0.05 and p < 0.0001 for viloxazine ER 100 mg and 200 mg vs placebo, respectively).</p>
- A third trial evaluated viloxazine ER in 310 patients aged 12 to 17 years of age who were randomized to viloxazine ER 200 mg, 400 mg, or placebo (*Nasser 2021a*). After 6 weeks of treatment, viloxazine ER 200 mg and 400 mg resulted in LS mean changes from baseline in ADHD-RS-5 total scores of -16.0, -16.5, and -11.4 for viloxazine ER 200 mg, 400 mg, and placebo, respectively (p < 0.05 vs placebo for both comparisons).</p>
- The Dyanavel XR (amphetamine ER) tablet was approved in November 2021 for the treatment of patients 6 years and older. The pharmacokinetic profile of the Dyanavel XR tablet was established to be bioequivalent to that of the Dyanavel XR oral suspension (*Dyanavel XR Prescribing Information 2022*). The safety and efficacy of the ER tablet were evaluated in a randomized, DB, PC, fixed-dose study in 130 adult patients with ADHD (*Cutler et al 2022a*). Patients entered a 5-week, DB, dose-titration phase in which they were randomized to receive Dyanavel XR tablet or matching placebo once daily in the morning. The starting dose of 5 mg was titrated in 5-mg increments per week, and patients received a final dose of 20 mg for 14 ± 3 days before visit 5. The primary endpoint of mean PERMP-Total score (PERMP-T) across all postdose time points at visit 5 was significantly higher (improved) in the Dyanavel XR tablet group compared to the placebo group (302.8 vs 279.6; p = 0.0043).
- Xelstrym (dextroamphetamine transdermal system) was approved by the FDA in 2022 for the treatment of ADHD in adults and pediatric patients aged 6 to 17 years. Its efficacy was supported by previous, well-controlled studies of lisdexamfetamine in pediatric and adult patients, in addition to a MC, DB, randomized, PC, modified analog classroom study in pediatric patients aged 6 to 17 years (*Cutler et al 2022b*). The study was conducted in 2 periods, and Xelstrym patches delivering different doses (5, 10, 15, and 20 mg) were evaluated. Patients were enrolled in a 5-week, OL, stepwise dose-optimization period in which they were started on a 5-mg patch and evaluated weekly for possible adjustments to the next dose level. Once the optimal dose was reached, it was maintained during a 2-week, crossover, DB treatment period. A total of 106 patients entered the DB treatment period. The study found Xelstrym to be associated with significant improvement in the SKAMP total score compared to placebo (LS mean difference -5.87; 95% CI, 6.76 to -4.97; p < 0.001).
- A systematic (Cochrane) review of 185 RCTs (*Storebø et al 2015*) (N = 12,245) in children and adolescents with ADHD found that methylphenidate may improve teacher-rated ADHD symptoms, teacher-reported general behavior, and parent-reported quality of life (QOL) vs placebo. However, the evidence was of low quality.
- An RCT called the Preschool ADHD Treatment Study (PATS) (Greenhill et al 2006) evaluated the efficacy of
 methylphenidate immediate-release (IR) in 303 preschool children with ADHD and found that it demonstrated
 significant reductions on ADHD symptom scales; however, the effect sizes (0.4 to 0.8) were smaller than those
 generally reported for school-age children.
- A systematic (Cochrane) review of 23 PC, RCTs (*Punja et al 2016*) (N = 2675) found that amphetamines were
 effective at improving the core symptoms of ADHD, but they were also associated with a higher risk of AEs compared
 to placebo. There was no evidence that one kind of amphetamine was better than another and there was no
 difference between short-acting and long-acting formulations.
- ∘ A meta-analysis of 25 DB, PC, RCTs (*Schwartz et al 2014*) (N = 3928) in children and adolescents with ADHD found atomoxetine to be superior to placebo for overall ADHD symptoms, with a medium effect size (-0.64).
- A meta-analysis of 25 RCTs (all rated as low or very low quality evidence) in children with autism and concurrent ADHD symptoms concluded that methylphenidate and atomoxetine both reduced parent-rated hyperactivity and inattention (*Rodrigues et al 2021*). Methylphenidate also reduced teacher-rated hyperactivity and inattention, but atomoxetine only reduced teacher-rated inattention.
- A meta-analysis of 12 RCTs (*Hirota et al 2014*) (N = 2276) in pediatric patients with ADHD found that alpha₂adrenergic agonists were significantly superior to placebo for overall ADHD symptoms both as monotherapy and, to a
 lesser extent, as augmentation therapy to stimulants.
 - Meta-analytic results failed to demonstrate a significant difference in efficacy between alpha₂-adrenergic agonists. In sub-analyses of individual formulations, the ER formulations separated robustly from placebo whereas the IR formulations did not separate from placebo.
- A systematic review of 16 RCTs and 1 meta-analysis (*Chan et al 2016*) (N = 2668) found evidence supporting the use of methylphenidate ER and amphetamine ER formulations, atomoxetine, and guanfacine ER for the treatment of ADHD in adolescents. For the primary outcome measure of mean change in ADHD-RS total symptom score, both stimulant and nonstimulant medications led to clinically significant reductions of 14.93 to 24.60 points.



- For the treatment of ADHD in children and adolescents, stimulants typically have a slightly larger treatment effect size (standardized mean difference [SMD]) than nonstimulants (approximately 1.0 vs approximately 0.7 for both atomoxetine and alpha₂-adrenergic agonists). However, there is insufficient evidence to definitively conclude that one stimulant is more efficacious than another (*Krull 2022e*, *Wolraich et al 2019*).
 - An Agency for Healthcare Research and Quality (AHRQ) review of 78 studies (*Jadad et al 1999*) evaluating the
 efficacy of various interventions for the treatment of ADHD in children and adults found few, if any, differences
 between methylphenidate and dextroamphetamine.
 - o A meta-analysis of 23 DB, PC trials (*Faraone 2010a*) comparing the efficacy of methylphenidate and amphetamine formulations found that amphetamine products may be moderately more efficacious than methylphenidate products.
 - o A DB, PC, RCT (*Newcorn et al 2008*) (N = 516) comparing the efficacy of atomoxetine vs methylphenidate ER (osmotic-release formulation) in patients 6 to 16 years of age with ADHD found that both drugs were superior to placebo in terms of response rate, and that methylphenidate ER was superior to atomoxetine.
 - A meta-analysis of 29 DB, PC trials (Faraone et al 2006) evaluated the efficacy of various medications (methylphenidate and amphetamine compounds, atomoxetine, pemoline [no longer available in the US], bupropion, and modafinil) for the treatment of ADHD. The effect sizes for nonstimulant medications were significantly less than those for IR stimulants or long-acting stimulants. The 2 classes of stimulant medications did not differ significantly from one another.
 - o A meta-analysis of 28 DB, PC, RCTs (*Stuhec et al 2015*) (N = 4699) compared the efficacy of various medications for the treatment of ADHD in children and adolescents. Efficacy in reducing ADHD symptoms compared to placebo was small for bupropion (SMD, -0.32; 95% CI, -0.69 to 0.05), modest for atomoxetine (SMD, -0.68; 95% CI, -0.76 to -0.59) and methylphenidate (SMD, -0.75; 95% CI, -0.98 to -0.52), and highest for lisdexamfetamine (SMD, -1.28; 95% CI, -1.84 to -0.71).
 - A network meta-analysis and mixed treatment comparison of 36 RCTs (*Joseph et al 2017*) evaluating the
 comparative efficacy and safety of ADHD pharmacotherapies in children and adolescents found that
 lisdexamfetamine had greater efficacy than guanfacine ER, atomoxetine, and methylphenidate ER. Guanfacine ER
 had a high posterior probability of being more efficacious than atomoxetine, but their credible intervals overlapped.
 - o A network meta-analysis of 48 DB, RCTs (*Padilha et al 2018*) compared the safety and efficacy of various ADHD medications in children and adolescents. Of the 12 trials that were evaluated for efficacy, analysis was performed using the Clinical Global Impression Improvement (CGI-I) scale for 3 drugs, which showed that methylphenidate was more effective than atomoxetine (MD, 3.15; 95% CI, 0.75 to 13.71) and guanfacine (MD, 1.92; 95% CI, 0.64 to 5.94). Thirty-three trials were evaluated for safety. Ranking of AEs showed that lisdexamfetamine was more likely to cause sleep disorders, loss of appetite, and behavior problems compared to other treatments.
- Alpha₂-adrenergic agonists have been associated with improvements in ADHD symptoms and comorbid tics.
 - A meta-analysis of 9 DB, PC, RCTs (*Bloch et al 2009*) (N = 477) was conducted to determine the relative efficacy of different medications in treating ADHD and tic symptoms in children with both Tourette syndrome and ADHD.
 - Methylphenidate seemed to offer the greatest improvement of ADHD symptoms and did not seem to worsen tic symptoms.
 - o Alpha₂-adrenergic agonists offered the best combined improvement in both tic and ADHD symptoms.
 - o Atomoxetine significantly improved both tic and ADHD severity compared to placebo.
 - o One small study found that tic severity was significantly increased with higher doses of dextroamphetamine treatment.
 - A Cochrane review of 8 RCTs (Osland et al 2018) including 510 children with both ADHD and a chronic tic disorder found low-quality evidence for improvement of ADHD symptoms with methylphenidate, atomoxetine, and clonidine, and very low-quality evidence for desipramine, dextroamphetamine, guanfacine, and deprenyl. Tic symptoms improved with guanfacine, desipramine, methylphenidate, clonidine, and a combination of methylphenidate and clonidine. The authors noted that in 1 study with a short duration (3 weeks), high doses of dextroamphetamine worsened tics
- There are limited efficacy data regarding the treatment of ADHD in the adult population. Comparison of effect sizes in clinical trials suggests that stimulant medications are more efficacious in adult ADHD than nonstimulants.
 - o In April 2022, the FDA approved an expanded indication for Qelbree for the treatment of ADHD in adults based on the results of a DB, MC, randomized, PC, flexible-dose, parallel-group monotherapy trial (*Qelbree Prescribing Information 2022, Nasser 2022*). A total of 374 patients with ADHD aged 18 to 65 years were randomized to receive viloxazine ER (flexible dose of 200 to 600 mg/day) or matching placebo for 6 weeks. The primary and secondary endpoints were the change in the Adult ADHD Investigator Symptom Rating Scale (AISRS) total score and the Clinical Global Impressions-Severity of Illness (CGI-S) score, respectively, from baseline at end of study. Patients in the viloxazine

Data as of October 11, 2022 HJ-U/KS-U/AVD

Page 162



ER group had a greater reduction in the AISRS total score than the placebo group (LS mean change, -15.5 vs -11.7; p = 0.0040). A significantly greater reduction in the CGI-S score was also seen in patients treated with viloxazine ER compared to placebo (LS mean change, -1.4 vs -1.0; p = 0.0023).

- In a meta-analysis of 12 clinical trials (*Cunill et al 2013*) (N = 3375) comparing atomoxetine with placebo in adult ADHD, atomoxetine led to a modestly greater reduction in ADHD symptom severity but was associated with higher all-cause discontinuation.
- A meta-analysis (Faraone 2010b) of 19 randomized trials of 13 medications for adult ADHD found a greater average
 effect size for reduction in ADHD symptoms in patients receiving short- and long-acting stimulant medications (vs
 placebo; 0.86 and 0.73, respectively) compared with patients receiving nonstimulant medication (vs placebo; 0.39).
 No difference in effect size was found between short- and long-acting stimulants.
- o A meta-analysis of 20 randomized trials (*Stuhec et al 2019*) compared the efficacy, acceptability, and tolerability of lisdexamfetamine, mixed amphetamine salts, methylphenidate, and modafinil in the treatment of ADHD in adults. The highest effect size in reducing ADHD symptoms was found with lisdexamfetamine (SMD -0.89; 95% CI, -1.09 to -0.70), while moderate reductions in symptoms were seen with mixed amphetamine salts (SMD -0.64; 95% CI, -0.83 to -0.45) and methylphenidate (SMD -0.50; 95% CI, -0.58 to -0.41). No efficacy was reported with modafinil.
- O A Cochrane review of 19 studies (Castells et al 2018, N = 2521) comparing dextroamphetamine, lisdexamfetamine, and mixed amphetamine salts for the treatment of ADHD in adults found that overall, amphetamines reduced the patient- and clinician-rated severity of ADHD symptoms compared to placebo; however, they did not improve retention in treatment. Amphetamines were associated with an increased proportion of patients who withdrew because of AEs. When comparing different types of amphetamines, lisdexamfetamine and mixed amphetamine salts reduced the severity of ADHD symptoms as rated by clinicians, but dextroamphetamine did not. No differences in any outcome were found when comparing immediate- and sustained-release formulations.
- o A systematic review and network meta-analysis (*Elliot et al 2020*) of 81 RCTs compared methylphenidate, atomoxetine, dexamfetamine, lisdexamfetamine, guanfacine, mixed amphetamine salts, modafinil, and bupropion for the treatment of ADHD in adults. Treatment with any ADHD pharmacotherapy was associated with statistically significant improvement in patient-reported clinical response vs placebo. When drugs were analyzed individually, only atomoxetine was found to significantly improve patient-reported clinical response compared to placebo (mean difference [MD], -5.9; 95% CI, -12.6 to -0.4). Atomoxetine (MD, -3.7; 95% CI, -6.7 to -0.9), sustained-release methylphenidate (MD, -5.7; 95% CI, -11.2 to -0.3), and low-dose methylphenidate (MD, -10.4; 95% CI, -19.0 to -2.1) were found to improve clinician-assessed clinical response compared to placebo. No significant differences were observed between individual medications when response was considered as a continuous outcome.
- Another meta-analysis (Cortese et al 2018) of 133 RCTs comparing the use of amphetamines, atomoxetine, bupropion, clonidine, guanfacine, methylphenidate, and modafinil for the treatment of ADHD found that all drugs were superior to placebo for ADHD core symptoms as rated by clinicians in children and adolescents, and all drugs except for modafinil were more efficacious than placebo in adults.
 - When comparing the various drugs based on teachers' ratings in children and adolescents, only methylphenidate and modafinil were found to be more efficacious than placebo.
 - In head-to-head comparisons, differences in efficacy based on clinicians' ratings were found, favoring amphetamines over modafinil (SMD, -0.39; 95% CI -0.67 to -0.12), atomoxetine (SMD, -0.46; 95% CI, -0.65 to -0.27), and methylphenidate (SMD, -0.24; 95% CI, -0.44 to -0.05) in children and adolescents. Efficacy results based on clinicians' ratings were similar for adults, and favored amphetamines over modafinil (SMD, -0.94; 95% CI -1.43 to -0.46), atomoxetine (SMD, -0.34; 95% CI, -0.58 to -0.10), and methylphenidate (SMD, -0.29; 95% CI, -0.54 to -0.05).
- Lisdexamfetamine dimesylate has demonstrated efficacy in the treatment of BED. Direct comparison trials between lisdexamfetamine and other drugs used off-label to treat BED are lacking.
 - o In 2 Phase 3, 12-week, randomized, DB, PC trials (*McElroy et al 2016*) (N = 773) in patients with moderate to severe BED, lisdexamfetamine-treated patients had a statistically significantly greater reduction from baseline in mean number of binge days per week at week 12 vs placebo (treatment difference in study 1: -1.35; 95% CI, -1.70 to -1.01; study 2: -1.66; 95% CI, -2.04 to -1.28; both p < 0.001).
 - A 12-month, OL extension study (*Gasior et al 2017*) (N = 599) in adults with BED found that the long-term safety and tolerability of lisdexamfetamine were generally consistent with the safety profile observed in 3 previous short-term trials in BED as well as its established profile for ADHD. Common treatment-emergent AEs included dry mouth, headache, insomnia, and upper respiratory tract infection. Weight loss and increases in blood pressure and pulse rate were also observed.



- o In a Phase 3, DB, randomized, PC, withdrawal study (*Hudson et al 2017*) (N = 418) in adults with moderate to severe BED, responders to lisdexamfetamine during a 12-week OL phase were randomized to placebo or continued lisdexamfetamine during a 26-week, DB phase. The percentage of patients meeting relapse criteria was 3.7% with lisdexamfetamine vs 32.1% with placebo; time to relapse statistically favored lisdexamfetamine (p < 0.001). The hazard ratio (HR) was 0.09 (95% CI, 0.04 to 0.23).
- A systematic review and meta-analysis of 14 clinical and 7 preclinical trials concluded that lisdexamfetamine effectively treats BED and reduces both symptoms (MD, 0.93; 95% CI, 0.74 to 1.12) and body weight (based on systematic review only) (*Schneider et al 2021*).
- o A systematic review and meta-analysis of 9 waitlist-controlled psychological trials and 25 PC trials evaluating pharmacologic (n = 19) or combination (n = 6) treatment for BED (*Brownley et al 2016*) found that therapist-led cognitive behavioral therapy (CBT), lisdexamfetamine, and second-generation antidepressants (SGAs) increased binge-eating abstinence (relative risk [RR], 4.95 [95% CI, 3.06 to 8.00], 2.61 [95% CI, 2.04 to 3.33], and 1.67 [95% CI, 1.24 to 2.26], respectively), while lisdexamfetamine and SGAs decreased binge-eating frequency (MD in days/week, -1.35 [95% CI, -1.77 to -0.93] and -0.67 [95% CI, -1.26 to -0.09], respectively). Topiramate and other forms of CBT also increased abstinence and reduced binge-eating frequency.
- A 2018 systematic review and meta-analysis of 45 RCTs (Ghaderi et al 2018) compared various psychological, pharmacological, and combined treatments for BED, and found moderate support for the efficacy of CBT and CBT-guided self-help (moderate quality of evidence), and low-quality evidence to support interpersonal psychotherapy, selective serotonin reuptake inhibitors (SSRIs), and lisdexamfetamine for the cessation of or reduction in the frequency of binge eating. Only lisdexamfetamine showed a modest effect on weight loss (SMD for body mass index 5.23; 95% CI, -6.52 to -3.94).

CLINICAL GUIDELINES

ADHD

- Several clinical guidelines have provided recommendations on the treatment of ADHD in children and adolescents.
 - o According to the American Academy of Pediatrics (AAP) guidelines (Wolraich et al 2019), the evidence is particularly strong for stimulant medications, and sufficient but less strong for atomoxetine, guanfacine ER, and clonidine ER (in that order; newer agents such as serdexmethylphenidate/dexmethylphenidate [Azstarys] and viloxazine [Qelbree] are not addressed in the current guidelines). Guanfacine ER and clonidine ER have evidence to support their use as adjunctive therapy with stimulant medications. Methylphenidate is recommended for preschool-aged children who have had an inadequate response to behavioral interventions.
 - o The Society for Developmental and Behavioral Pediatrics guideline on assessment and treatment of children and adolescents with complex ADHD states that treatment should aim to improve functional impairment and include skill development in self-management strategies (*Barbaresi et al 2020*). Multimodal treatment with both behavioral and pharmacologic therapies may be needed. Specific pharmacologic classes are discussed in the context of learning disorder, for which the guideline recommends both stimulants and atomoxetine, with stimulants having a greater strength of evidence, and autism, for which a stimulant is recommended first followed by an alpha₂-adrenergic agonist or atomoxetine. Stimulant use is also endorsed in children with intellectual disability, tics, anxiety or depression, and disruptive behavior disorders.
 - The Medical Letter recommends that treatment of ADHD in school-age children or adults should begin with a stimulant, either a methylphenidate- or amphetamine-based formulation (*Med Lett Drugs Ther 2020*). Mixing short- and long-acting stimulants can be helpful to achieve an immediate effect for early-morning school classes or for reducing rebound irritability or overactivity, especially in the evening. Nonstimulants can be used in combination with stimulants or when stimulants are contraindicated, ineffective, or not tolerated.
 - o According to the American Academy of Neurology guidelines for treatment of tics (*Pringsheim et al 2019*), physicians should counsel individuals with tics and comorbid ADHD that alpha₂-adrenergic agonists may provide benefit for both conditions. Alpha₂-adrenergic agonists and topiramate should be prescribed for the treatment of tics when the benefits of treatment outweigh the risks, while antipsychotics and botulinum toxin may be prescribed when the benefits outweigh the risks.
 - The American Academy of Child and Adolescent Psychiatry (AACAP) practice parameter for the treatment of children
 and adolescents with tic disorders (*Murphy et al 2013*) states that alpha₂-adrenergic agonists have demonstrated an
 effect size of 0.5 for the amelioration of tics and may be preferred by some prescribers over antipsychotics due to
 their relatively favorable AE profile.



Narcolepsy

• The American Academy of Sleep Medicine (AASM) practice parameters (*Maski et al 2021*) recommend various drugs for the treatment of daytime sleepiness in adults due to narcolepsy including modafinil, pitolisant, sodium oxybate, solriamfetol (strongly recommended), and armodafinil, dextroamphetamine, and methylphenidate (conditionally recommended). Idiopathic hypersomnia in adults should be treated with modafinil (strongly recommended), clarithromycin, methylphenidate, pitolisant, or sodium oxybate (conditionally recommended). Recommended therapies for children with narcolepsy include modafinil and sodium oxybate (both conditionally recommended).

BED

- According to the American Psychiatric Association (APA) practice guidelines on eating disorders (Yager et al 2006, Yager et al 2012 [guideline watch update], now categorized as a legacy guideline), treatment of BED may include the following:
 - Nutritional rehabilitation and counseling
 - Psychosocial treatment
 - CBT, behavior therapy, dialectical behavior therapy (DBT), and interpersonal therapy (IPT) have all been associated with binge frequency reduction rates of 67% or more and significant abstinence rates during active treatment.
 - Self-help programs using self-guided, professionally designed manuals have been effective in reducing the symptoms of BED in the short-run for some patients and may have long-term benefit.
 - Medications
 - Antidepressant treatment is associated with short-term reductions in binge-eating but generally does not result in substantial weight loss. SSRIs have the fewest difficulties with AEs and the most evidence for efficacy when used at the high end of the recommended dose range.
 - Topiramate can reduce bingeing and decrease weight, but its use may be limited by AEs.
 - Combination psychotherapy and pharmacotherapy
 - For most patients, adding antidepressant therapy to a behavioral weight control and/or CBT regimen does not have a significant effect on binge suppression.
 - Although limited evidence is available, combined treatment is frequently used in clinical practice.
- The American Association of Clinical Endocrinologists and the American College of Endocrinology (AACE/ACE) guidelines for medical care of patients with obesity (*Garvey et al 2016*) recommend the following for patients with overweight or obesity who have BED:
 - Patients should be treated with a structured behavioral/lifestyle program, combined with CBT or other psychological interventions
 - Treatment with orlistat or approved medications containing topiramate or bupropion may be considered in conjunction with structured lifestyle therapy, CBT, and/or psychological interventions
- The Task Force on Eating Disorders of the World Federation of Societies of Biological Psychiatry (*Aigner et al 2011*) concluded that for the treatment of BED, grade A evidence supports the use of imipramine (moderate risk-benefit ratio), sertraline (good risk-benefit ratio), citalopram/escitalopram (good risk-benefit ratio), orlistat (low to moderate risk-benefit ratio), and topiramate (moderate risk-benefit ratio). Atomoxetine has grade B evidence supporting its use.

SAFETY SUMMARY

- Due to the potential for abuse, the stimulants are classified as Schedule II controlled substances. Atomoxetine, clonidine ER, quanfacine ER, and viloxazine ER are not classified as controlled substances.
- Various stimulants are contraindicated for use in patients with advanced arteriosclerosis, symptomatic CV disease, moderate to severe hypertension, hyperthyroidism, hypersensitivity to sympathomimetic amines, glaucoma, agitated states, history of drug abuse, tics, and in those using monoamine oxidase inhibitors (MAOIs). The stimulants carry a boxed warning for potential drug abuse and dependence. They also have warnings for increased risks of serious CV reactions, psychiatric AEs, suppression of growth, seizures, visual disturbance, peripheral vasculopathy, and priapism. Amphetamines have a warning for risk of serotonin syndrome when used in combination with other drugs affecting the serotonergic neurotransmitter systems.
 - o Common AEs of stimulants include anorexia, decreased weight, tachycardia, anxiety, irritability, and insomnia.
 - Refer to the prescribing information for details on warnings, precautions, and AEs for individual products. For example:
 - QuilliChew ER can be harmful to patients with phenylketonuria (PKU) since it contains phenylalanine.



- Because Concerta and Relexxii tablets are nondeformable and do not appreciably change in shape in the gastrointestinal tract, they should not ordinarily be administered to patients with preexisting severe gastrointestinal narrowing.
- The use of Daytrana and Xelstrym may lead to contact sensitization; in addition, exposure of the application site to external heat sources should be avoided due to increased absorption of the drug. Daytrana use may result in chemical leuokoderma.
- Adhansia XR capsules contain FD&C yellow No. 5 dye (tartrazine), which may cause allergic-type reactions in susceptible patients.
- Atomoxetine is contraindicated for use in patients with narrow angle glaucoma, pheochromocytoma, severe CV disorders, hypersensitivity to any component of the product, and in those taking MAOIs. It carries a boxed warning for a rare increased risk of suicidal ideation in children and adolescents. It also has warnings for serious CV events, effects on blood pressure and heart rate, effects on growth, psychotic or manic symptoms, aggressive behavior or hostility, rare cases of severe liver injury, urinary retention, and priapism. Patients should be screened for a personal or family history of bipolar disorder prior to use of atomoxetine due to the risk of activation of mania or hypomania.
 - o Common AEs associated with atomoxetine include somnolence, nausea, and vomiting.
- Viloxazine ER is contraindicated with concurrent use of MAOIs and sensitive CYP1A2 substrates or CYP1A2 substrates
 with a narrow therapeutic index. Viloxazine ER carries a boxed warning for suicidal thoughts and behavior in patients
 treated with the drug. It also has warnings for effects on heart rate and blood pressure and the potential for somnolence
 and fatigue. Patients should be screened for bipolar disorder prior to use of viloxazine ER due to the risk of activation of
 mania or hypomania.
 - o Common AEs associated with viloxazine ER include somnolence, nausea, and vomiting.
- The alpha₂-adrenergic agonists are contraindicated in patients known to be hypersensitive to any constituent of the product. They carry warnings for increased risk of hypotension, bradycardia, and syncope; sedation and somnolence; rebound hypertension; and cardiac conduction abnormalities.
 - o Common AEs associated with clonidine ER include somnolence, fatigue, and irritability while common AEs with guanfacine ER include somnolence, fatigue, and hypotension.

DOSING AND ADMINISTRATION

Table 4. Dosing and Administration

Drug	Duration of action*	Available Formulations	Route	Usual Recommended Frequency	Comments
Stimulants					
Evekeo (amphetamine)	4 to 6 h	Tablets	Oral	ADHD, narcolepsy: Daily up to divided doses daily Exogenous obesity: Divided doses daily	ADHD and narcolepsy The first dose should be given upon awakening; additional doses at intervals of 4 to 6 hours.
Evekeo ODT (amphetamine)	4 to 6 h	Orally disintegrating tablets	Oral	Once or twice daily in the morning	As soon as the blister pack is opened, the tablet should be placed on the patient's tongue and allowed to disintegrate without chewing or crushing. The tablet will disintegrate in saliva so that it can be swallowed.

Data as of October 11, 2022 HJ-U/KS-U/AVD

Page 166

This information is considered confidential and proprietary to OptumRx. It is intended for internal use only and should be disseminated only to authorized recipients. The contents of the therapeutic class overviews on this website ("Content") are for informational purposes only. The Content is not intended to be a substitute for professional medical advice, diagnosis, or treatment. Patients should always seek the advice of a physician or other qualified health provider with any questions regarding a medical condition. Clinicians should refer to the full prescribing information and published resources when making medical decisions.



Drug	Duration of action*	Available Formulations	Route	Usual Recommended Frequency	Comments
Adzenys XR-ODT (amphetamine ER)	10 to 12 h	Orally disintegrating tablets	Oral	Daily in the morning	As soon as the blister pack is opened, the tablet should be placed on the patient's tongue and allowed to disintegrate without chewing or crushing. The tablet will disintegrate in saliva so that it can be swallowed.
Dyanavel XR (amphetamine ER)	Up to 13 h	Suspension, ER tablets	Oral	Daily in the morning	The bottle should be shaken before administration. ER tablets may be chewed or swallowed whole. The 5 mg tablet may be split along the score line.
Adderall (mixed amphetamine salts)	4 to 6 h	Tablets	Oral	ADHD, narcolepsy: Daily up to divided doses daily	The first dose should be given on awakening, then additional doses at intervals of 4 to 6 hours.
Adderall XR (mixed amphetamine salts ER)	10 to 12 h	Capsules	Oral	Daily in the morning	Capsules may be taken whole, or the capsule may be opened and the entire contents sprinkled on applesauce and consumed immediately. The dose of a single capsule should not be divided.
Mydayis (mixed amphetamine salts ER)	16 h	Capsules	Oral	Daily in the morning	Dosage adjustment is needed for severe renal impairment. Use in end stage renal disease (ESRD) is not recommended. Capsules may be taken whole, or the



Drug	Duration of action*	Available Formulations	Route	Usual Recommended Frequency	Comments
					capsule may be opened and the entire contents sprinkled on applesauce and consumed immediately in its entirety without chewing. The dose of a single capsule should not be divided.
Focalin (dexmethylphenidate)	3 to 5 h	Tablets	Oral	Twice daily	Separate doses by at least 4 hours.
Focalin XR (dexmethylphenidate ER)	8 to 12 h	Capsules	Oral	Daily in the morning	ER capsules may be taken whole, or the capsule may be opened and the entire contents sprinkled on applesauce and consumed immediately in its entirety without chewing. The dose of a single capsule should not be divided.
ProCentra, Zenzedi (dextroamphetamine)	4 to 6 h	Solution (ProCentra) Tablets (Zenzedi)	Oral	ADHD, narcolepsy: Daily up to divided doses daily	The first dose should be given upon awakening; additional doses at intervals of 4 to 6 hours
Dexedrine Spansule (dextroamphetamine SR)	6 to 8 h	Capsules	Oral	ADHD Daily or twice daily Narcolepsy Daily	
Xelstrym (dextroamphetamine transdermal system)	Up to 12 h	Transdermal system	Transdermal	The patch should be applied 2 hours before an effect is needed and removed within 9 hours.	Dose titration and final dosage should be individualized depending on clinical response and tolerability. Dosage adjustment is needed for renal impairment/ESRD.



Drug	Duration of action*	Available Formulations	Route	Usual Recommended Frequency	Comments
Vyvanse (lisdexamfetamine)	10 to 12 h	Capsules, chewable tablets	Oral	ADHD, BED: Daily in the morning	Dosage adjustment is needed for renal impairment/ESRD. The capsules may be swallowed whole or can be opened, emptied, and mixed with yogurt, water, or orange juice and consumed immediately. A single capsule should not be divided. The chewable tablets must be chewed thoroughly before swallowing. A single dose should not be divided.
Desoxyn (methamphetamine)	4 to 5 h	Tablets	Oral	Daily to twice daily	
Methylin, Ritalin (methylphenidate)	3 to 5 h	Chewable tablets, tablets (Ritalin), solution (Methylin)	Oral		The chewable tablets should be taken with at least 8 ounces (a full glass) of water or other fluid. The liquid and chewable tablets
Methylphenidate ER	8 h	Tablets		Twice daily to 3 times daily	should be given 30 to 45 minutes before meals. The ER tablets may be used in place of the IR tablets when the 8-hour dosage of the ER product corresponds to the titrated 8-hour dosage of the IR products. The ER tablets must be swallowed



Drug	Duration of action*	Available Formulations	Route	Usual Recommended Frequency	Comments
					whole and never crushed or chewed.
Adhansia XR (methylphenidate ER)	13 h	Capsules	Oral	Daily in the morning	The capsules may be taken whole or they can be opened and sprinkled onto applesauce or yogurt; the entire contents of the mixture should be consumed within 10 minutes, and should not be chewed. The dose of a single capsule should not be divided.
Aptensio XR (methylphenidate ER)	12 h	Capsules	Oral	Daily in the morning	The capsules may be taken whole or they can be opened and sprinkled onto applesauce; the applesauce should be consumed immediately and it should not be chewed. The dose of a single capsule should not be divided.



Drug	Duration of action*	Available Formulations	Route	Usual Recommended Frequency	Comments
Concerta (methylphenidate ER) Methylphenidate ER	12 h	Tablets	Oral	Daily in the morning	The tablets should not be chewed or crushed. Note: An FDA analysis of methylphenidate ER products manufactured by UCB/Kremers (formerly Kudco) and Mallinckrodt indicated that in some individuals, they may deliver the drug in the body at a slower rate during the 7- to 12-hour range. As a result, the FDA changed the therapeutic equivalence of these products from AB to BX. Because these manufacturers have subsequently failed to demonstrate that their products are bioequivalent to the brand-name reference drug, the FDA proposed to withdraw their approval (FDA 2016).
Cotempla XR-ODT (methylphenidate ER)	12 h	Orally disintegrating tablets	Oral	Daily in the morning	As soon as the blister pack is opened, the tablet should be placed on the patient's tongue and allowed to disintegrate without chewing or crushing. The tablet will disintegrate in saliva so that it can be swallowed.



Drug	Duration of action*	Available Formulations	Route	Usual Recommended Frequency	Comments
Jornay PM (methylphenidate ER)	10 h	Capsules	Oral	Daily in the evening	The capsule may be swallowed whole or it may be opened and the contents sprinkled onto applesauce and given immediately. The capsule contents must not be crushed or chewed, the dose of a single capsule should not be divided, and the contents of the entire capsule should be taken at the same time.
Methylphenidate ER (CD)	6 to 9 h	Capsules	Oral	Daily in the morning	The capsule may be swallowed whole or it may be opened and the contents sprinkled onto a small amount (one tablespoon) of applesauce and given immediately, followed by some fluids. The capsule contents must not be crushed or chewed.
QuilliChew ER (methylphenidate ER)	8 h	Chewable tablets	Oral	Daily in the morning	A 10 mg or 15 mg dose can be achieved by breaking in half the functionally scored 20 mg and 30 mg tablets, respectively.
Quillivant XR (methylphenidate ER)	12 h	Suspension	Oral	Daily in the morning	The bottle of Quillivant XR should be shaken vigorously for 10 seconds prior to administration. The suspension is stable for up to 4



Drug	Duration of action*	Available Formulations	Route	Usual Recommended	Comments
		1 official actions		Frequency	months once
Relexxii (methylphenidate ER 72 mg)	12 h	Tablet	Oral	Daily in the morning	reconstituted. The tablet must be swallowed whole with the aid of liquids, and must not be chewed, divided, or crushed.
Ritalin LA (methylphenidate ER)	6 to 9 h	Capsules	Oral	Daily in the morning	The capsule may be swallowed whole or may be administered by sprinkling the capsule contents on a small amount of applesauce; the contents should not be crushed, chewed, or divided. The mixture should be consumed immediately.
Daytrana (methylphenidate transdermal system)	Up to 12 h	Transdermal system	Transdermal	The patch should be applied 2 hours before an effect is needed and removed within 9 hours. It may be removed earlier than 9 hours if a shorter duration of effect is desired or late day side effects appear.	
Azstarys (serdexmethylphenidate/ dexmethylphenidate)	10 to 13 h	Capsules	Oral	Daily in the morning	The capsule may be swallowed whole or may be administered by sprinkling the capsule contents over 2 tablespoons of applesauce or 50 mL of water. The mixture should be consumed immediately.
Non-stimulants	At least 10			Daily in the	Dosage adjustment
Strattera (atomoxetine)	to 12 h	Capsules	Oral	morning or	is recommended for



Drug	Duration of action*	Available Formulations	Route	Usual Recommended	Comments
				frequency divided dose in the morning and late afternoon/ early evening	patients with moderate or severe hepatic insufficiency, for use with strong CYP2D6 inhibitors, and for patients known to be CYP2D6 poor metabolizers. The capsules are not intended to be opened and should be taken whole
Kapvay (clonidine ER)	At least 10 to 12 h	Tablets	Oral	Daily at bedtime or twice daily divided doses	be taken whole. With twice daily dosing, either an equal or higher split dosage should be given at bedtime. The tablets should not be crushed, chewed, or broken prior to swallowing. The initial dosage should be based on the degree of renal impairment.
Intuniv (guanfacine ER)	At least 8 to 12 h	Tablets	Oral	Daily in the morning or evening	The tablets should not be crushed, chewed, or broken prior to swallowing; they should not be administered with high fat meals, due to increased exposure. It may be necessary to reduce the dosage in patients with significant renal and hepatic impairment.
Qelbree (viloxazine ER)	Throughout the day (specific duration	Capsules	Oral	Daily	The capsule may be swallowed whole or may be administered by



Drug	Duration of action*	Available Formulations	Route	Usual Recommended Frequency	Comments
	not reported)				sprinkling the capsule contents over a teaspoon of applesauce. The mixture should be consumed within 2 hours, without chewing.

See the current prescribing information for full details

*References: Prescribing information for individual products, Medical Letter 2020, Pharmacist's Letter 2021, Krull 2022d.

CONCLUSION

- Both CNS stimulants and nonstimulants may be used for the treatment of ADHD. In general, stimulants are first-line treatment due to their superior efficacy. Clinical evidence suggests that methylphenidate and amphetamines are equally efficacious, but some patients may respond to one stimulant and not the other. Various short-, intermediate- and long-acting formulations (eg, tablets/capsules, chewable/orally disintegrating tablets, solution/suspension, transdermal patch) are available to provide a range of dosing options. Although nonstimulants such as atomoxetine and alpha2-adrenergic agonists have smaller effect sizes, they may be used in patients who have failed or are intolerant to stimulants or when there is concern about possible abuse or diversion. The efficacy of the nonstimulant viloxazine ER in comparison to other nonstimulants is unknown. The alpha2-adrenergic agonists are approved both as monotherapy and as adjunctive therapy to stimulants, and they have been shown to improve both tic and ADHD symptoms in patients with comorbid tic disorder.
 - Current consensus clinical guidelines for the treatment of children and adolescents with ADHD recommend that stimulants are highly effective for reducing core symptoms of ADHD in children (Wolraich et al 2019).
- Ultimately, the choice of the initial agent for treatment of ADHD depends upon various factors such as: duration of desired coverage; ability of the child to swallow pills; coexisting tic disorder (use of alpha₂-adrenergic agonists may be warranted); potential AEs, history of substance abuse in the patient or household member (eg, avoid stimulants or use stimulants with less potential for abuse [eg, lisdexamfetamine, osmotic-release preparation, methylphenidate patch]); and preference of the patient and parent/guardian (*Krull 2022b*).
- Various stimulants are indicated for treatment of narcolepsy and are generally considered to be second-line agents after modafinil/armodafinil due to their sympathomimetic AEs (*Scammell 2021*).
- Lisdexamfetamine is the only FDA-approved drug indicated for the treatment of moderate to severe BED, with demonstrated efficacy in reduction of mean binge days per week vs placebo. Direct comparison trials between lisdexamfetamine and other drugs used off-label to treat BED are lacking.

REFERENCES

- Adderall [package insert], Parsippany, NJ: Teva Pharmaceuticals USA, Inc.; February 2022.
- Adderall XR [package insert], Lexington, MA: Takeda Pharmaceuticals America, Inc.; February 2022.
- Adhansia XR FDA clinical review. Food and Drug Administration website. January 29, 2019.
 https://www.accessdata.fda.gov/drugsatfda docs/nda/2019/212038Orig1s000MedR.pdf. Accessed October 10, 2022.
- Adhansia XR [package insert], Wilson, NC: Purdue Pharmaceuticals L.P.; June 2021.
- Adzenys XR-ODT [package insert], Grand Prairie, TX: Neos Therapeutics, Inc.; March 2022.
- Aigner M, Treasure J, Kaye W, Kasper S; WFSBP Task Force on Eating Disorders. World Federation of Societies of Biological Psychiatry (WFSBP) guidelines for the pharmacological treatment of eating disorders. World J Biol Psychiatry. 2011;12(6):400-443.
- Aptensio XR [package insert], Coventry, RI: Rhodes Pharmaceuticals, L.P.; June 2021.
- Azstarys [package insert], Grand Rapids, MI: Corium, Inc.; June 2021.
- Barbaresi WJ, Campbell L, Diekroger EA, et al. Society for Developmental and Behavioral Pediatrics clinical practice guideline for the assessment and treatment of children and adolescents with complex attention-deficit/hyperactivity disorder. *J Dev Behav Pediatr.* 2020;41 Suppl 2S:S35-S57. doi:10.1097/DBP.00000000000000770
- Bloch MH, Panza KE, Landeros-Weisenberger A, Leckman JF. Meta-analysis: treatment of attention-deficit/hyperactivity disorder in children with comorbid tic disorders. J Am Acad Child Adolesc Psychiatry. 2009;48(9):884-893. doi:10.1097/CHI.0b013e3181b26e9f



- Brownley KA, Berkman ND, Peat CM, et al. Binge-eating disorder in adults: A systematic review and meta-analysis. Ann Intern Med. 2016;165(6):409-420
- Bukstein O. Attention deficit hyperactivity disorder in adults: Epidemiology, pathogenesis, clinical features, course, assessment, and diagnosis.
 UpToDate Web site. 2022. http://www.uptodate.com. Updated April 7, 2022. Accessed October 4, 2022.
- Castells X, Blanco-Silvente L, Cunill R. Amphetamines for attention deficit hyperactivity disorder (ADHD) in adults. Cochrane Database Syst Rev. 2018;8:CD007813. doi: 10.1002/14651858.CD007813.pub3.
- Centers for Disease Control and Prevention (CDC). Attention Deficit Hyperactivity Disorder (ADHD). National Center for Health Statistics. https://www.cdc.gov/nchs/fastats/adhd.htm.

 Reviewed June 11, 2021. Accessed October 4, 2022.
- Chan E, Fogler JM, Hammerness PG. Treatment of attention-deficit/hyperactivity disorder in adolescents: a systematic review. JAMA. 2016;315(18):1997-2008.
- Childress AC, Brams M, Cutler AJ, et al. The efficacy and safety of Evekeo, racemic amphetamine sulfate, for treatment of attentiondeficit/hyperactivity disorder symptoms: a multicenter, dose-optimized, double-blind, randomized, placebo-controlled crossover laboratory classroom study. J Child Adolesc Psychopharmacol. 2015;25(5):402-414. doi: 10.1089/cap.2014.0176
- Childress AC, Cutler AJ, Marraffino A, et al. A randomized, double-blind, placebo-controlled study of HLD200, a delayed-release and extended-release
 methylphenidate, in children with attention-deficit/hyperactivity disorder: An evaluation of safety and efficacy throughout the day and across settings. J
 Child Adolesc Psychopharmacol. 2020;30(1):2-14.
- Childress AC, Kollins SH, Cutler AJ, Marraffino A, Sikes CR. Efficacy, safety, and tolerability of an extended-release orally disintegrating
 methylphenidate tablet in children 6-12 years of age with attention-deficit/hyperactivity disorder in the laboratory classroom setting. *J Child Adolesc Psychopharmacol*. 2017;27(1):66-74.
- Clinical Pharmacology Website. https://www.clinicalkey.com/pharmacology/login. Accessed October 4, 2022.
- Concerta [package insert], Titusville, NJ: Janssen Pharmaceuticals, Inc.; April 2022.
- Cotempla XR-ODT [package insert], Grand Prairie, TX: Neos Therapeutics Brands, LCC; June 2021.
- Cortese S, Adamo N, Del Giovane C, et al. Comparative efficacy and tolerability of medications for attention-deficit hyperactivity disorder in children, adolescents, and adults: a systematic review and network meta-analysis. *Lancet Psychiatry*. 2018;5(9):727-738. doi: 10.1016/S2215-0366(18)30269-4.
- Cunill R, Castells X, Tobias A, Capellà D. Atomoxetine for attention deficit hyperactivity disorder in the adulthood: a meta-analysis and meta-regression. Pharmacoepidemiol Drug Saf. 2013;22(9):961-969.
- Cutler AJ, Childress AC, Pardo A, et al. Randomized, double-blind, placebo-controlled, fixed-dose study to evaluate the efficacy and safety of amphetamine extended-release tablets in adults with attention-deficit/hyperactivity disorder. J Clin Psychiatry. 2022a;83(5):22m14438. doi:10.4088/JCP.22m14438
- Cutler AJ, Suzuki K, Starling B, et al. Efficacy and safety of dextroamphetamine transdermal system for the treatment of attention-deficit/hyperactivity disorder in children and adolescents: results from a pivotal phase 2 study. J Child Adolesc Psychopharmacol. 2022b;32(2):89-97. doi:10.1089/cap.2021.0107
- Daytrana [package insert], Miami, FL: Noven Therapeutics, LLC; June 2021.
- Desoxyn [package insert], Lebanon, NJ: Recordati Rare Diseases Inc.; March 2019.
- Dexedrine Spansule [package insert], Bridgewater, NJ: Amneal Pharmaceuticals LLC; January 2022.
- Drugs@FDA [database on the Internet], Rockville (MD): Food and Drug Administration (US), Center for Drug Evaluation and Research; 2022.
 Available from: http://www.accessdata.fda.gov/scripts/cder/daf/index.cfm. Accessed October 4, 2022.
- Drugs for ADHD. Med Lett Drugs Ther. 2020;62 (1590):9-16.
- Dyanavel XR [package insert], Monmouth Junction: Tris Pharma, Inc.; May 2022.
- Elliott J, Johnston A, Husereau D, et al. Pharmacologic treatment of attention deficit hyperactivity disorder in adults: A systematic review and network meta-analysis. *PLoS One*. 2020;15(10):e0240584. doi: 10.1371/journal.pone.0240584
- Evekeo [package insert], Atlanta, GA: Arbor Pharmaceuticals, LLC; October 2019.
- Evekeo ODT [package insert], Atlanta, GA: Arbor Pharmaceuticals, LLC; April 2021.
- Faraone SV, Biederman J, Spencer TJ, Aleardi M. Comparing the efficacy of medications for ADHD using meta-analysis. MedGenMed. 2006;8(4):4.
- Faraone SV, Buitelaar J. Comparing the efficacy of stimulants for ADHD in children and adolescents using meta-analysis. *Eur Child Adolesc Psychiatry*, 2010a;19(4):353-364.
- Faraone SV, Glatt SJ. A comparison of the efficacy of medications for adult attention-deficit/hyperactivity disorder using meta-analysis of effect sizes. *J Clin Psychiatry*. 2010b;71(6):754-763.
- Feldman HM, Reiff MI. Clinical practice. Attention deficit-hyperactivity disorder in children and adolescents. *N Engl J Med*. 2014;370(9):838-846.
- Focalin [package insert], East Hanover, NJ: Novartis Pharmaceuticals Corporation.; June 2021.
- Focalin XR [package insert], East Hanover, NJ: Novartis Pharmaceuticals Corporation.; June 2021.
- Food and Drug Administration. Methylphenidate Hydrochloride Extended Release Tablets (generic Concerta) made by Mallinckrodt and Kudco. FDA
 Web site. https://www.fda.gov/drugs/drug-safety-and-availability/methylphenidate-hydrochloride-extended-release-tablets-generic-concerta-made-mallinckrodt-and-kudco. Updated November 4, 2016. Accessed October 4, 2022.
- Frick G, Yan B, Adler LA. Triple-bead mixed amphetamine salts (SHP465) in adults with ADHD: results of a phase 3, double-blind, randomized, forced-dose trial. *J Atten Disord*. 2020;24(3):402-413. doi: 10.1177/1087054717696771
- Garvey WT, Mechanick JI, Brett EM, et al; Reviewers of the AACE/ACE Obesity Clinical Practice Guidelines. American Association of Clinical Endocrinologists and American College of Endocrinology comprehensive clinical practice guidelines for medical care of patients with obesity. *Endocr Pract*. 2016;22 Suppl 3:1-203. doi: 10.4158/EP161365.GL.
- Gasior M, Hudson J, Quintero J, Ferreira-Cornwell MC, Radewonuk J, McElroy SL. A phase 3, multicenter, open-label, 12-month extension safety and tolerability trial of lisdexamfetamine dimesylate in adults with binge eating disorder. *J Clin Psychopharmacol*. 2017;37(3):315-322.
- Ghaderi A, Odeberg J, Gustafsson S, et al. Psychological, pharmacological, and combined treatments for binge eating disorder: a systematic review and meta-analysis. PeerJ. 2018;6:e5113. doi:10.7717/peerj.5113.
- Greenhill L, Kollins S, Abikoff H, et al. Efficacy and safety of immediate-release methylphenidate treatment for preschoolers with ADHD. J Am Acad Child Adolesc Psychiatry. 2006;45(11):1284-1293.



- Hirota T, Schwartz S, Correll CU. Alpha-2 agonists for attention-deficit/hyperactivity disorder in youth: a systematic review and meta-analysis of monotherapy and add-on trials to stimulant therapy. J Am Acad Child Adolesc Psychiatry. 2014;53(2):153-173.
- Intuniv [package insert], Lexington, MA: Shire US Inc.; August 2020.
- Hudson JI, McElroy SL, Ferreira-Cornwell MC, Radewonuk J, Gasior M. Efficacy of lisdexamfetamine in adults with moderate to severe binge-eating disorder: A randomized clinical trial. *JAMA Psychiatry*. 2017;74(9):903-910.
- Jadad AR, Boyle M, Cunningham C, Kim M, Schachar R. Treatment of attention-deficit/hyperactivity disorder. Evid Rep Technol Assess (Summ).
 1999:(11):i-viii, 1-341.
- Jornay PM [package insert], Cherry Hill, NJ: Ironshore Pharmaceuticals, Inc.; June 2021.
- Joseph A, Ayyagari R, Xie M, et al. Comparative efficacy and safety of attention-deficit/hyperactivity disorder pharmacotherapies, including guanfacine extended release: a mixed treatment comparison. Eur Child Adolesc Psychiatry. 2017;26(8):875-897.
- Kapvay [package insert], St. Michael, Barbados: Concordia Pharmaceuticals, Inc.: February 2020.
- Kollins SH, Braeckman R, Guenther S, et al. A Randomized, controlled laboratory classroom dtudy of serdexmethylphenidate and d-methylphenidate capsules in children with attention-deficit/hyperactivity disorder. *J Child Adolesc Psychopharmacol*. 2021;31(9):597-609.
- Krull KR. Attention deficit hyperactivity disorder in children and adolescents: Clinical features and diagnosis. UpToDate Web site. 2022a. http://www.uptodate.com. Updated May 31, 2022. Accessed October 4, 2022.
- Krull KR. Attention deficit hyperactivity disorder in children and adolescents: Epidemiology and pathogenesis. UpToDate Web site. 2022b. http://www.uptodate.com. Updated May 16, 2022. Accessed October 4, 2022.
- Krull KR. Attention deficit hyperactivity disorder in children and adolescents: Overview of treatment and prognosis. UpToDate Web site. 2022d. http://www.uptodate.com. Updated June 6, 2022. Accessed October 4, 2022.
- Krull KR. Attention deficit hyperactivity disorder in children and adolescents: Treatment with medications. UpToDate Web site. 2022e. http://www.uptodate.com. Updated August 10, 2022. Accessed October 4, 2022.
- Krull KR. Pharmacology of drugs used to treat attention deficit hyperactivity disorder in children and adolescents. UpToDate Web site. 2022c. http://www.uptodate.com. Updated August 10, 2022. Accessed October 4, 2022.
- Maski K, Trotti LM, Kotagal S, et al. Treatment of central disorders of hypersomnolence: an American Academy of Sleep Medicine clinical practice guideline. J Clin Sleep Med. 2021;17(9):1881-1893. doi:10.5664/jcsm.9328
- McElroy SL, Hudson J, Ferreira-Cornwell MC, Radewonuk J, Whitaker T, Gasior M. Lisdexamfetamine dimesylate for adults with moderate to severe binge eating disorder: Results of two pivotal phase 3 randomized controlled trials. *Neuropsychopharmacology*. 2016;41(5):1251-1260.
- Methylin oral solution [package insert], Florham Park, NJ: Shionogi Inc.; July 2021.
- Methylphenidate chewable tablets [package insert], Central Islip, NY: Ascent Pharmaceuticals, Inc.; August 2021.
- Methylphenidate ER [package insert], Newtown, PA: KVK-Tech, Inc.; April 2021.
- Methylphenidate ER (CD) [package insert], Philadelphia, PA: Lannett Company, Inc.; May 2021.
- Methylphenidate hydrochloride extended-release tablets (generic Concerta) made by Mallinckrodt and Kudco. FDA Web site. November 4, 2016.
- Murphy TK, Lewin AB, Storch EA, Stock S; American Academy of Child and Adolescent Psychiatry (AACAP) Committee on Quality Issues (CQI).
 Practice parameter for the assessment and treatment of children and adolescents with tic disorders. J Am Acad Child Adolesc Psychiatry.
 2013;52(12):1341-1359.
- Mydayis [package insert], Lexington, MA: Shire US Inc.; January 2022.
- Nasser A, Hull JT, Chaturvedi SA, et al. A phase III, randomized, double-blind, placebo-controlled trial assessing the efficacy and safety of viloxazine extended-release capsules in adults with attention-deficit/hyperactivity disorder. CNS Drugs. 2022;36(8):897-915. doi:10.1007/s40263-022-00938-w
- Nasser A, Liranso T, Adewole T, et al. A Phase III, randomized, placebo-controlled trial to assess the efficacy and safety of once-daily SPN-812 (viloxazine extended-release) in the treatment of attention-deficit/hyperactivity disorder in school-age children. Clin Ther. 2020;42(8):1452-1466.
- Nasser A, Liranso T, Adewole T, et al. A phase 3, placebo-controlled trial of once-daily viloxazine extended-release capsules in adolescents with attention-deficit/hyperactivity disorder. J Clin Psychopharmacol. 2021a;41(4):370-380. doi:10.1097/JCP.000000000001404
- Nasser A, Liranso T, Adewole T, et al. Once-daily SPN-812 200 and 400 mg in the treatment of ADHD in school-aged children: A Phase III
 randomized. controlled trial. Clin Ther. 2021b;S0149-2918(21)00054-0.
- Newcorn JH, Kratochvil CJ, Allen AJ, et al. Atomoxetine/Methylphenidate Comparative Study Group. Atomoxetine and osmotically released
 methylphenidate for the treatment of attention deficit hyperactivity disorder: acute comparison and differential response. *Am J Psychiatry*.
 2008;165(6):721-730.
- Orange Book: Approved Drug Products with Therapeutic Equivalence Evaluations. Food and Drug Administration Web site. http://www.accessdata.fda.gov/scripts/cder/ob/default.cfm. Accessed October 4, 2022.
- Osland ST, Steeves TD, Pringsheim T. Pharmacological treatment for attention deficit hyperactivity disorder (ADHD) in children with comorbid tic disorders. Cochrane Database Syst Rev. 2018;6:CD007990. doi:10.1002/14651858.CD007990.pub3.
- Padilha SCOS, Virtuoso S, Tonin FS, Borba HHL, Pontarolo R. Efficacy and safety of drugs for attention deficit hyperactivity disorder in children and adolescents: a network meta-analysis. *Eur Child Adolesc Psychiatry*. 2018;27(10):1335-1345. doi: 10.1007/s00787-018-1125-0.
- PL Detail-Document, Comparison of ADHD medications. Pharmacist's Letter/Prescriber's Letter. September 2021.
- Pliszka SR, Wilens TE, Bostrom S, et al. Efficacy and safety of HLD200, delayed-release and extended-release methylphenidate, in children with attention-deficit/hyperactivity disorder. J Child Adolesc Psychopharmacol. 2017;27(6):474-482. doi: 10.1089/cap.2017.0084.
- Practice parameter for the assessment and treatment of children and adolescents with tic disorders. J Am Acad Child Adolesc Psychiatry. 2013;52(12):1341-1359.
- Pringsheim T, Okun MS, Müller-Vahl K, et al. Practice guideline recommendations summary: Treatment of tics in people with Tourette syndrome and chronic tic disorders. *Neurology*. 2019;92(19):896-906.
- ProCentra [package insert], Newport, KY: Independence Pharmaceuticals, LLC; December 2021.
- Punja S, Shamseer L, Hartling L, et al. Amphetamines for attention deficit hyperactivity disorder (ADHD) in children and adolescents. Cochrane Database Syst Rev. 2016;2:CD009996.
- Qelbree [package insert], Rockville, MD: Supernus Pharmaceuticals, Inc.; April 2022.
- QuilliChew ER [package insert], Monmouth Junction, NJ: Tris Pharma, Inc.; June 2021.

Data as of October 11, 2022 HJ-U/KS-U/AVD



- Quillivant XR [package insert], Monmouth Junction, NJ: Tris Pharma, Inc.; June 2021.
- Relexxii [package insert], Alpharetta, GA: Vertical Pharmaceuticals, LLC; November 2021.
- Rodrigues R, Lai MC, Beswick A, et al. Practitioner Review: Pharmacological treatment of attention-deficit/hyperactivity disorder symptoms in children
 and youth with autism spectrum disorder: a systematic review and meta-analysis. *J Child Psychol Psychiatry*. 2021;62(6):680-700.
 doi:10.1111/jcpp.13305
- Ritalin [package insert], East Hanover, NJ: Novartis Pharmaceuticals Corporation; June 2021.
- Ritalin LA [package insert], East Hanover, NJ: Novartis Pharmaceuticals Corporation.; June 2021.
- Scammell TE. Treatment of narcolepsy in adults. UpToDate Web site. 2022. http://www.uptodate.com. Updated September 2, 2022. Accessed October 4, 2022.
- Schneider E, Higgs S, Dourish CT. Lisdexamfetamine and binge-eating disorder: A systematic review and meta-analysis of the preclinical and clinical data with a focus on mechanism of drug action in treating the disorder. *Eur Neuropsychopharmacol*. 2021;53:49-78. doi:10.1016/j.euroneuro.2021.08.001
- Schwartz S, Correll CU. Efficacy and safety of atomoxetine in children and adolescents with attention-deficit/hyperactivity disorder: results from a comprehensive meta-analysis and metaregression. J Am Acad Child Adolesc Psychiatry. 2014;53(2):174-187.
- Storebø OJ, Ramstad E, Krogh HB, et al. Methylphenidate for children and adolescents with attention deficit hyperactivity disorder (ADHD). Cochrane Database Syst Rev. 2015;11:CD009885.
- Strattera [package insert], Indianapolis, IN: Lilly USA, Inc.; January 2022.
- Stuhec M, Lukić P, Locatelli I. Efficacy, acceptability, and tolerability of lisdexamfetamine, mixed amphetamine salts, methylphenidate, and modafinil in the treatment of attention-deficit hyperactivity disorder in adults: a systematic review and meta-analysis. *Ann Pharmacother*. 2019;53(2):121-133. doi:10.1177/1060028018795703.
- Stuhec M, Munda B, Svab V, Locatelli I. Comparative efficacy and acceptability of atomoxetine, lisdexamfetamine, bupropion and methylphenidate in treatment of attention deficit hyperactivity disorder in children and adolescents: a meta-analysis with focus on bupropion. *J Affect Disord*. 2015;178:149-159.
- Vyvanse [package insert], Lexington, MA: Takeda Pharmaceuticals America, Inc.; January 2022.
- Weisler RH, Greenbaum M, Arnold V, et al. Efficacy and safety of SHP465 mixed amphetamine salts in the treatment of attention-deficit/hyperactivity disorder in adults: results of a randomized, double-blind, placebo-controlled, forced-dose clinical study. CNS Drugs. 2017;31(8):685-697.
- Wigal T, Brams M, Frick G, Yan B, Madhoo M. A randomized, double-blind study of SHP465 mixed amphetamine salts extended-release in adults with ADHD using a simulated adult workplace design. *Postgrad Med.* 2018a;130(5):481-493. doi: 10.1080/00325481.2018.1481712
- Wigal T, Childress A, Frick G, Yan B, Wigal S, Madhoo M. Effects of SHP465 mixed amphetamine salts in adults with ADHD in a simulated adult workplace environment. *Postgrad Med.* 2018b;130(1):111-121. doi: 10.1080/00325481.2018.1389227
- Wigal T, Lopez F, Frick G, Yan B, Robertson B, Madhoo M. A randomized, double-blind, 3-way crossover, analog classroom study of SHP465 mixed amphetamine salts extended-release in adolescents with ADHD. Postgrad Med. 2019;131(3):212-224. doi: 10.1080/00325481.2019.1574402
- Wigal SB, Wigal T, Childress A, Donnelly GAE, Reiz JL. The time course of effect of multilayer-release methylphenidate hydrochloride capsules: a randomized, double-blind study of adults with ADHD in a simulated adult workplace environment. *J Atten Disord*. 2020;24(3):373-383. doi: 10.1177/1087054716672335
- Wolraich ML, Hagan JF Jr., Allan C, et al; Subcommittee on Children and Adolescents with Attention-Deficit/Hyperactive Disorder. Clinical practice guideline for the diagnosis, evaluation, and treatment of attention-deficit/hyperactivity disorder in children and adolescents. *Pediatrics*. 2019;144(4):e20192528. doi: 10.1542/peds.2019-2528
- Yager J, Devlin MF, Halmi KA, et al. Guideline watch (August 2012): Practice guideline for the treatment of patients with eating disorders, 3rd edition. Psychiatry Online Web site. http://psychiatryonline.org/guidelines. Accessed October 4, 2022.
- Yager J, Devlin MF, Halmi KA, et al. Practice guideline for the treatment of patients with eating disorders, 3rd edition (2006). Psychiatry Online Web site. http://psychiatryonline.org/quidelines. Accessed October 4, 2022.
- Zenzedi [package insert], Atlanta, GA: Arbor Pharmaceuticals, LLC; January 2022.
- Xelstrym [package insert], Miami, FL: Noven Pharmaceuticals, Inc.; March 2022.

Publication Date: October 31, 2022