The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

ADD/ADHD/ NARCOLEPSY AGENTS	Prior authorization (PA) is required for ADD/ADHD/Narcolepsy agents for patients 21 years of age or older. PA is also required for all non-preferred agents, regardless of age, the first day of therapy. Payment for a non-preferred agent will be authorized only for cases in which there is documentation of previous trial and therapy failure with a preferred agent, unless evidence is provided that use of these agents would be medically contraindicated. *If a non-preferred long-acting medication is requested, a trial if the preferred immediate release and extended release product of the same chemical entity is required, unless evidence is provided that use of these products would be medically contraindicated. Prior approval shall be granted if there is documentation of one of the following:
Use ADD/ADHD/Narcolepsy Agents PA form	 Attention deficit disorder. Attention deficit hyperactivity disorder. Narcolepsy. Other FDA approved indications
Alpha ₂ Agonists,	Prior authorization is required for extended-release alpha ₂ agonists. Payment will be considered for patients when the
Extended-Release	following is met:
Intuniv [™] Kapvay [™] Use Alpha ₂ Agonists, Extended-Release PA form	 The patient has a diagnosis of ADHD and is between 6 and 17 years of age; and Previous trial with the preferred immediate release product of the same chemical entity at a therapeutic dose that resulted in a partial response with a documented intolerance; and Previous trial and therapy failure at a therapeutic dose with one preferred amphetamine and one preferred non-amphetamine stimulant; and Previous trial and therapy failure at a therapeutic dose with atomoxetine (Strattera®). The required trials may be overridden when documented evidence is provided that the use of these agents would be medically contraindicated.
Alpha ₁ Proteinase	Prior authorization is required for Alpha ₁ -Proteinase Inhibitor enzymes. Payment will be authorized only for cases in which
Inhibitor Enzymes	there is a diagnosis of congenital alpha ₁ -proteinase inhibitor (alpha ₁ -PI; alpha1-antitrypsin) deficiency with clinically
Use Miscellaneous PA form Amylino Mimetic	demonstrable panacinar emphysema. Payment for a non-preferred Alpha ₁ -Proteinase Inhibitor enzyme will be authorized only for cases in which there is documentation of previous trial and therapy failure with a preferred agent. Prior authorization is required for amylino mimetics (Symlin®). Payment will be considered under the following
(Symlin®) Use Amylino Mimetic (Symlin®) form	conditions: 1) Diagnosis of Type 1 or Type 2 diabetes mellitus, 2) Concurrent use of insulin therapy, 3) Documentation of blood glucose monitoring three or more times daily, 4) Inadequate reduction in HbgA1C despite multiple titration with basal/bolus insulin dosing regiments. Initial authorizations will be approved for six months; additional prior authorizations will be considered on an individual basis after review of medical necessity and documented improvement in HbgA1C since the beginning of the initial prior authorization period.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Angiotensin Receptor	Payment for Angiotensin Receptor Blockers (ARB) and Angiotensin Receptor Blocker Combinations will only be
Blocker Before ACE	considered for cases in which there is a contraindication or therapy failure with at least one ACE-I or ACE-I Combination.
Inhibitor	A completed prior authorization form will need to be submitted if a trial with an ACE-I or ACE-I Combination of at least
	30 days in length is not found in the point-of-sale system and/or unless evidence is provided that use of an ACE-I or ACE-I
	Combination would be medically contraindicated. Prior authorization is required for all non-preferred ARBs and ARB
	Combinations the first day of therapy. Payment for a non-preferred ARB or ARB Combination will be considered
Use Angiotensin Receptor Blocker Before ACE	following documentation of recent trials and therapy failures with a preferred ACE-I or ACE-I Combination AND a
Inhibitor PA form	preferred ARB or ARB Combination.
Anti-Acne	Prior authorization is required for all prescription topical acne products. Payment for the treatment of mild to moderate acne
	vulgaris will be considered under the following conditions:
	1. Previous trial and therapy failure with a preferred over-the-counter benzoyl peroxide product, which is covered by
	the program without prior authorization.
	2. Payment for non-preferred topical acne products will be authorized only for cases in which there is documentation
	of previous trials and therapy failures with two preferred topical agents of a different chemical entity.
	3. If the patient presents with a preponderance of comedonal acne, topical retinoid products may be utilized as first
	line agents with prior authorization (use Topical Retinoids PA form).
	4. Requests for non-preferred combination products may only be considered after documented separate trials and
	therapy failures with the individual ingredients.
Use Anti-Acne PA form	The required trials may be overridden when documented evidence is provided that the use of these agents would be
	medically contraindicated.
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The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

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Antiemetic-5HT3	Prior authorization is required for preferred Antiemetic-5H	T3 Receptor Antagonists/Substance P Neurokinin medications
Receptor	for quantities exceeding the following dosage limits per mo	onth. Payment for Antiemetic-5HT3 Receptor Agonists/
Antagonists/	Substance P Neurokinin Agents beyond this limit will be considered on an individual basis after review of submitted	
Substance P	documentation.	
Neurokinin Agents	Prior authorization will be required for all non-preferred Antiemetic-5HT3 Receptor Antagonists/Substance P Neurokinin	
0		r non-preferred medications will be authorized only for cases in
		py failure with a preferred agent in this class. Note: Aprepitant
	(Emend®) will only be payable when used in combination	
	dexamethasone) for patients receiving highly emetogenic c	
	Aprepitant (N)/Emend® (P):	Ondansetron (P)/Zofran® (N):
	Aprepliant (N) Emend (1). $4 - 125$ mg capsules	12 – 4mg tablets
	8 – 80mg capsules	12 – 4mg tablets 12 – 8mg tablets
	Dolasetron (N)/Anzemet® (N):	4 – 24mg tablets
	5 - 50 mg/100 mg tablets	50mL/month – oral solution (4mg/5mL)
	4 vials (100mg/5mL)	4 – 20mL vials (2mg/mL)
	8 ampules (12.5mg/0.625mL)	8 – 2mL vials (2mg/mL)
	Granisetron (N)/Kytril [®] /Granisol [™] (N):	Ondansetron ODT (P)/Zofran® ODT (N):
	8 – 1mg tablets	12 – 4mg tablets
	30mL – oral solution (1mg/5mL)	12 – 8mg tablets
Use Antiemetic-5HT3	8 vials (1mg/mL)	Palonosetron (N)/ Aloxi [®] (N):
Receptor Antagonists/	2 vials (4mg/mL)	4 vials (0.25mg/5mL)
Substance P Neurokinin		
Agents form		
Anti-Fungal		ngal therapy for a cumulative 90 days of therapy per 12-month
		all non-preferred oral antifungal therapy beginning the first day
	of therapy. Payment for a non-preferred oral antifungal wil	l be authorized only for cases in which there is documentation of
	previous trial and therapy failure with a preferred agent. Pa	yment for any oral antifungal therapy beyond a cumulative 90
	days of therapy per 12-month period per patient will be aut	horized in cases where the patient has a diagnosis of an
		tion. This prior authorization requirement does not apply to
Use Anti-Fungal PA form	nystatin.	1 11 7
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The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

-	Updated 7/30/2012
Antihistamines	Prior authorization is required for all non-preferred antihistamines and preferred 2 nd generation prescription antihistamines.
	Patients 21 years of age and older must have three unsuccessful trials with antihistamines that do not require prior authorization, prior to the approval of a non-preferred 1 st generation or preferred 2 nd generation prescription antihistamine. Two of the trials must be with cetirizine and loratadine. Prior to approval of a non-preferred 2 nd generation antihistamine, in addition to the above criteria, there must be an unsuccessful trial with a preferred 2 nd generation prescription antihistamine.
	Patients 20 years of age and younger must have unsuccessful trials with cetirizine and loratadine prior to the approval of a non-preferred 1 st generation or preferred 2 nd generation prescription antihistamine. Prior to approval of a non-preferred 2 nd generation antihistamine, in addition to the above criteria, there must be an unsuccessful trial with a preferred 2 nd generation prescription antihistamine.
Use Antihistamine PA form	The required trials may be overridden when documented evidence is provided that the use of these agents would be medically contraindicated.
Anti-Thrombotics, Injectable	Prior authorization is required for use of any preferred injectable anti-thrombotic agent longer than 10 consecutive days. Prior authorization will be required for all non-preferred injectable anti-thrombotic agents beginning the first day of therapy. Payment for non-preferred anti-thrombotic injectable agents will be authorized only for cases in which there is documentation of previous trial and therapy failure with a preferred agent. Payment for usage of injectable anti-thrombotic agents beyond this limit will be authorized for cases in which there is a clinical diagnosis of: 1. Pregnancy or planned pregnancy 2. Cancer-associated thromboembolic disease 3. Anti-thrombin III deficiency
Use Anti-Thrombotic Injectable PA form	4. Warfarin allergy5. History of thrombotic event while on therapeutic anticoagulant therapy.6. Total hip arthroplasty.
Becaplermin	Prior authorization is required for Regranex [®] . Payment for new prescriptions will be authorized for ten weeks for patients
(Regranex [®])	who meet the following criteria:
	1. Diagnosis of lower extremity diabetic neuropathic ulcers that extend into the subcutaneous tissue or beyond
	2. Inadequate response to 2 weeks of wound debridement and topical moist wound dressing Longer than 10 weeks will be authorized for patients who meet the following criteria:
Use Regranex® PA form	Wound has decreased in size by 30% after 10 weeks
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The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

	Updated 7/30/2012
Benzodiazepines	Prior authorization is required for non-preferred benzodiazepines. Payment for non-preferred benzodiazepines will be authorized in cases with documentation of previous trial and therapy failure with two preferred products. Requests for clobazam (ONFI) will be considered for a diagnosis of seizures associated with Lennox-Gastaut syndrome (LGS) in patients 2 years of age and older when used as an adjunctive treatment. Prior authorization will be approved for up to 12 months for documented: 1. Generalized anxiety disorder. 2. Panic attack with or without agoraphobia. 3. Seizure. 4. Non-progressive motor disorder. 5. Dystonia.
Use Benzodiazepine PA form	If a long-acting medication is requested, one of the therapeutic trials must include the immediate release form of the requested benzodiazepine. Prior authorization requests will be approved for up to a three-month period for all other diagnoses related to the use of benzodiazepines. The required trials may be overridden when documented evidence is provided that use of these agents would be medically
	contraindicated.
Biologicals for Ankylosing Spondylitis Adalimumab (Humira®) Etanercept (Enbret®) Infliximab (Remicade®) Golimumab (Simponi™) Use Biologicals for Ankylosing Spondylitis PA form	Prior authorization is required for biologicals used for ankylosing spondylitis. Payment will be considered following inadequate responses to at least two preferred non-steroidal anti-inflammatories (NSAIDs) at maximum therapeutic doses, unless there are documented adverse responses or contraindications to NSAID use. These trials should be at least three months in duration. Patients with symptoms of peripheral arthritis must also have failed a 30-day treatment trial with at least one conventional disease modifying antirheumatic drug (DMARD), unless there is a documented adverse response or contraindication to DMARD use. DMARDs include sulfasalazine and methotrexate. Payment for non-preferred biologicals for ankylosing spondylitis will be considered only for cases in which there is documentation of previous trials and therapy failures with two preferred biological agents.
Biologicals for Arthritis Abatacept (Orencia®) Adalimumab (Humira®) Anakinra (Kineret®) Certolizumab Pegol (Cimzia®) Etanercept (Enbret®) Infliximab (Remicade®) Golimumab (Simponi™)	Prior authorization is required for biologicals used for arthritis. Payment will be considered following an inadequate response to two preferred disease modifying antirheumatic drugs (DMARD) used in combination. The combination must include methotrexate plus another preferred oral DMARD (hydroxycholoroquine, sulfasalazine, leflunomide, or minocycline). The required trials may be overridden when documented evidence is provided that the use of these agents would be medically contraindicated. Upon an unsuccessful methotrexate trial, the combination trial with a second DMARD may be overridden if there is evidence of severe disease documented by radiographic erosions. Payment for non-preferred biologicals for arthritis will be considered only for cases in which there is documentation of a previous trials and therapy failures with two preferred biological agents.
Use Biologicals for Arthritis PA form	

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

Biologicals for Inflammatory Bowel Disease	Prior authorization is required for biologicals used for inflammatory bowel disease. Payment for non-preferred biologicals for inflammatory bowel disease will be considered only for cases in which there is documentation of a previous trial and therapy failure with a preferred agent.
Adalimumab (Humira [®]) Certolizumab Pegol (Cimzia [®]) Infliximab (Remicade [®])	 Crohn's Disease – Payment will be considered following an inadequate response to two preferred conventional therapy including aminosalicylates (mesalamine, sulfasalazine), azathioprine/6-mercaptopurine, and/or methotrexate. Ulcerative colitis (moderate to severe) – Payment will be considered following an inadequate response to two preferred conventional therapies including aminosalicylates and azathioprine/6-mercaptopurine.
Use Biologicals for Inflammatory Bowel Disease PA form	
Biologicals for	Prior authorization is required for biologicals used for plaque psoriasis. Payment will be considered following an inadequate
Plaque Psoriasis Alefacept (Amevive®) Adalimumab (Humira®) Etanercept (Enbret®) Infliximab (Remicade®) Use Biologicals for Plaque Psoriasis PA form	response to phototherapy, systemic retinoids (oral isotretinoin), methotrexate, or cyclosporine. Payment for non-preferred biologicals for plaque psoriasis will be considered only for cases in which there is documentation of previous trials and therapy failures with two preferred biological agents.
Buprenorphine (Butrans [™]) Transdermal System	 Prior authorization is required for Butrans[™]. Payment will be considered when the following conditions are met: 1. Previous trials and therapy failures at a therapeutic dose with long acting morphine sulfate product and methadone. The preferred trials must allow for adequate dose titration and show use of a short acting narcotic for breakthrough pain. 2. A trial and therapy failure with fentanyl patch at maximum tolerated dose.
Use Buprenorphine (Butrans™) Transdermal System PA form	The required trials may be overridden when documented evidence it provided that use of these agents would be medically contraindicated.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

Chronic Pain Syndromes

Duloxetine (Cymbalta[®]) Pregabalin (Lyrica[®]) Milnacipran (Savella[™]) A prior authorization is required for duloxetine (Cymbalta[®]), pregabalin (Lyrica[®]), and milnacipran (Savella^{TM}). Payment will be considered under the following conditions:

- 1. A diagnosis of fibromyalgia (Cymbalta[®], Lyrica[®], and Savella[™])
 - a. a trial and therapy failure at a therapeutic dose with three drugs from three distinct therapeutic classes from the following: tricyclic antidepressant, muscle relaxant, SSRI/SNRI, tramadol, or gabapentin, **WITH**
 - b. documented non-pharmacologic therapies (cognitive behavior therapies, exercise, etc.), AND
 - c. documentation of a previous trial and therapy failure at a therapeutic dose with Savella $^{\text{\tiny TM}}$ when Cymbalta $^{\text{\tiny RM}}$ and Lyrica $^{\text{\tiny RM}}$ are requested.
- 2. A diagnosis of post-herpetic neuralgia (Lyrica®)

A trial and therapy failure at a therapeutic dose with at least two drugs from two distinct therapeutic classes from the following: tricyclic antidepressant, topical lidocaine, valproate, carbamazepine, or gabapentin.

3. A diagnosis of diabetic peripheral neuropathy (Cymbalta® and Lyrica®)

A trial and therapy failure at a therapeutic dose with at least two drugs from two distinct therapeutic classes from the following: tricyclic antidepressant, topical lidocaine, tramadol, or gabapentin.

- 4. A diagnosis of partial onset seizures, as adjunct therapy (Lyrica®)
- 5. A diagnosis of major depressive disorder or generalized anxiety disorder (Cymbalta[®])
- 6. A diagnosis of chronic musculoskeletal pain (Cymbalta®)

A trial and therapy failure at a therapeutic dose with at least three drugs from three distinct therapeutic classes from the following: NSAIDs, opioids, tramadol, or tricyclic antidepressants.

Use Chronic Pain Syndromes PA form

Requests for concomitant use of these agents for an indicated chronic pain diagnosis may only be considered once each agent has been tried at maximum tolerated dose separately. Duplicate use of drugs from the same therapeutic category will not be considered.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

Colchicine (Colcrys®)	Prior authorization is not required for colchicine (Colcrys®) for the treatment of acute gout for three (3) tablets per 60-day
	period. Prior authorization is required for colchicine (Colcrys®) for the treatment of chronic hyperuricemia/gout
	prophylaxis or Familial Mediterranean fever. Payment will be considered under the following conditions:
	1. Chronic hyperuricemia/gout prophylaxis following a trial and therapy failure at a therapeutic dose with allopurinol or
	probenecid. A quantity limit of sixty (60) tablets per thirty (30) days will be applied, when criteria for coverage are
	met.
	2. Familial Mediterranean fever. A maximum quantity of 120 tablets per thirty (30) days will be applied for this
	diagnosis.
Use Colchicine (Colcrys®)	The required trials may be overridden when documented evidence is provided that the use of these agents would be
PA form	medically contraindicated.
G TITE	
Concurrent IM/PO	A prior authorization is required for concurrent long acting injectable and oral antipsychotic medications after 12 weeks
Antipsychotic Use	(84 days) of concomitant treatment. Consideration of concomitant therapy beyond 12 weeks (84 days) will require
	documentation of medical necessity. Prior authorization is required for all non-preferred antipsychotics as indicated on the
	Iowa Medicaid Preferred Drug List beginning the first day of therapy. Payment for
Use Concurrent IM/PO	non-preferred antipsychotics will be considered only for cases in which there is documentation of previous trials and
Antipsychotic Utilization PA form	therapy failures with a preferred agent.
John	
Crizotinib (Xalkori®)	Prior authorization is required for Xalkori® (crizotinib). Payment will be considered for patients when the following is met:
	1. Diagnosis of locally advanced or metastatic non-small cell lung cancer (NSCLC) that is anaplastic lymphoma kinase
	(ALK)-positive as detected by an FDA-approved test (attach copy of results); and
Use Xalkori® PA form	2. Is prescribed by an oncologist.
DPP-4 Inhibitors	Prior authorization is required for dipeptidyl peptidase-4 (DPP-4) inhibitors and DPP-4 Inhibitor Combinations. Payment
	will be considered under the following conditions:
	1. A diagnosis of Type 2 Diabetes Mellitus
	2. Patient is 18 years of age or older
	3. The patient has not achieved HbgA1C goals using a combination of two or more antidiabetic medications (metformin,
	sulfonylurea, thiazolidinedione, or insulin) at maximum tolerated doses unless otherwise contraindicated.
	, , , , , , , , , , , , , , , , , , , ,
	Initial authorizations will be approved for six months; additional prior authorizations will be considered on an individual
	basis after review of medical necessity and documented improvement in HbgA1C since the beginning of the initial prior
	authorization period. Payment for a non-preferred agent will be authorized only for cases in which there is documentation
Use DPP-4 Inhibitor PA	of previous trial and therapy failure with a preferred agent, unless evidence is provided that use of these agents would be
form	medically contraindicated.
	medicany contamidence.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

	Opulated 7/50/2012
Dalfampridine	Prior authorization is required for dalfampridine (Ampyra [™]). Payment will be considered under the following conditions:
(Ampyra TM)	1. For patients that have a gait disorder associated with MS.
	2. Initial authorizations will be approved for 12 weeks with a baseline Timed 25-foot Walk (T25FW) assessment.
	3. Additional prior authorizations will be considered at 6 month intervals after assessing the benefit to the patient as
	measured by a 20% improvement in the T25FW from baseline. Renewal will not be approved if the 20% improvement
	is not maintained.
Use Dalfampridine	Prior authorizations will not be considered for patients with a seizure diagnosis or in patients will moderate to severe renal
(Ampyra TM) PA form	impairment.
Dextromethorphan	Prior authorization is required for Nuedexta TM . Payment will be considered under the following conditions:
and Quinidine	1. Patients must have a diagnosis of pseudobulbar affect (PBA) secondary to amyotrophic lateral sclerosis (ALS) or
(Nuedexta TM)	multiple sclerosis (MS).
(Nucucata)	2. A trial and therapy failure at a therapeutic dose with amitriptyline or an SSRI.
	3. Initial authorizations will be approved for 12 weeks with a baseline Center for Neurologic Studies Lability Scale
	(CNS-LS) questionnaire.
Use Dextromethorphan and	4. Subsequent prior authorizations will be considered at 6 month intervals with documented efficacy as seen in an
Quinidine (Nuedexta [™]) PA form	improvement in the CNS-LS questionnaire.
3	Discount of the form of the state of the sta
Digestive Enzymes	Prior authorization is required for all digestive enzymes. Payment for preferred digestive enzymes will be authorized only
	for cases in which there is a clinical diagnosis of malabsorption due to pancreatic insufficiency. Payment for non-
	preferred digestive enzymes will be authorized only for cases in which there is documentation of previous trials and
Use Miscellaneous PA form	therapy failures with two preferred products.
Dornase Alfa	Prior authorization is required for Pulmozyme [®] . Payment will be authorized only for cases in which there is a diagnosis of
(Pulmozyme®)	cystic fibrosis.
Use Miscellaneous PA form	
Duloxetine	See Chronic Pain Syndromes Prior Authorization Criteria.
(Cymbalta®)	
Use Chronic Pain Syndromes PA form	
Eplerenone	Prior authorization is required for Inspra [®] . Payment will be authorized only in cases where there is documented trial and
(Inspra [®])	therapy failure on Aldactone® or documented cases of gynecomastia from Aldactone® therapy.
(Inspra)	therapy famule on Aldactone of documented cases of gynecomastia from Aldactone therapy.
Use Miscellaneous PA form	

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

	Optiated 1/30/2012
Erythropoiesis	Prior authorization is required for erythropoiesis stimulating agents prescribed for outpatients for the treatment of anemia.
Stimulating Agents	Payment for non-preferred erythropoiesis stimulating agents will be authorized only for cases in which there is
	documentation of previous trial and therapy failure with a preferred agent.
	Patients who meet all of the following criteria may receive prior authorization for the use of erythropoiesis stimulating
	agents:
	1. Hemoglobin less than 10g/dL.If renewal of prior authorization is being requested, a hemoglobin less than 11g/dL (or
	less than 10g/dL for patients with Chronic Kidney Disease (CKD) not on dialysis) will be required for continued
	treatment. Hemoglobin laboratory values must be dated within four weeks of the prior authorization request.
	2. Transferrin saturation greater than or equal to 20 percent (transferrin saturation is calculated by dividing serum iron by
	the total iron binding capacity), ferritin levels greater than or equal to 100 mg/ml, or on concurrent therapeutic iron
	therapy. Transferrin saturation or ferritin levels must be dated within three months of the prior authorization request.
	3. For HIV-infected patients, the endogenous serum erythropoietin level must be less than or equal to 500 mU/ml to
Use Erythropoesis	initiate therapy.
Stimulating Agent PA form	4. No evidence of untreated GI bleeding, hemolysis, or Vitamin B-12, iron or folate deficiency.
Extended Release	Payment for a non-preferred extended release formulation will be considered when the following criteria are met:
Formulations	1. Previous trial and therapy failure with the preferred immediate release product of the same chemical entity at a
	therapeutic dose that resulted in a partial response with a documented intolerance and
	2. Previous trial and therapy failure at a therapeutic dose with a preferred drug of a different chemical entity
	indicated to treat the submitted diagnosis.
	The required trials may be overridden when documented evidence is provided that use of these agents would be medically
	contraindicated.
	contramateacea.
	Prior authorization is required for the following extended release formulation(s):
	Adoxa, Amrix, Augmentin XR, Cardura XL, Cipro XR, Coreg CR, Doryx, Flagyl ER, glipizide er, Glucotrol XL,
	Keppra XR, Lamictal XR, Lescol XL, Luvox CR, metronidazole sr, Mirapex ER, Moxatag, Paxil CR, Prozac Weekly,
Use Extended Release	Requip XL, Ryzolt, Seroquel XR, Solodyn ER, Tegretol XR, tramadol sr, Ultram ER.
Formulations PA form	
Febuxostat (Uloric®)	Prior authorization is required for febuxostat (Uloric®). Payment for febuxostat (Uloric®) will only be considered for cases
W. E. I. (III. I R) D.	in which symptoms of gout still persist while currently using 300mg per day of a preferred allopurinol product unless
Use Febuxostat (Uloric®) PA form	documentation is provided that such a trial would be medically contraindicated.
Fentanyl, Short	Prior authorization is required for short acting oral fentanyl products. Payment will be considered only if the diagnosis is
Acting Oral Products	for breakthrough cancer pain in opioid tolerant patients. These products carry a Black Box Warning .
Acting Oral Froducts	Actiq [®] , Fentora [®] , & Onsolis [™] :
	Are indicated only for the management of breakthrough cancer pain in patients with malignancies already
	receiving and tolerant to opioid therapy for their underlying persistent cancer pain.
Use Short Acting Oral	
Fentanyl Products PA form	Are contraindicated in the management of acute or postoperative pain. Because life-threatening hypoventilation and a sound account action and action and action actions are actions as a sound account action account action account action account action account a
	could occur at any dose in patients not taking chronic opiates, do not use in opioid non-tolerant patients.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Fifteen Day Initial	Designated drugs are limited to a fifteen day initial supply. These drugs are identified on the Fifteen Day Initial
Prescription Supply	Prescription Supply Limit list located on the website www.iowamedicaidpdl.com under the Preferred Drug Lists tab.
Limit	Providers must submit a prior authorization request for override consideration. Documentation of medical necessity,
	excluding patient convenience, is required for consideration of the fifteen day initial supply override.
Use Fifteen Day Initial	
Prescription Supply Limit	
PA form	
Fingolimod	Prior authorization is required for fingolimod (Gilenya [™]). Payment will be considered under the following conditions:
(Gilenya [™])	1. A diagnosis of relapsing forms of multiple sclerosis, and
	2. A previous trial and therapy failure with a preferred interferon and non-interferon used to treat multiple sclerosis.
	The required trial may be overridden when documented evidence is provided that the use of these agents would be
	medically contraindicated.
Granulocyte Colony	Prior authorization is required for therapy with granulocyte colony stimulating factor agents. Payment for non-preferred
Stimulating Factor	granulocyte colony stimulating factor agents will be authorized only for cases in which there is documentation of previous
Agents	trial and therapy failure with a preferred agent. Laboratory values for complete blood and platelet count must be contained
	as directed by the manufacturer's instructions. Dosage reduction and discontinuation of therapy may be required based on
	the manufacturer's guidelines. Payment shall be authorized for one of the following uses:
	1. Prevention or treatment of febrile neutropenia in patients with malignancies who are receiving myelosuppressive
	anticancer therapy.
	2. Treatment of neutropenia in patients with malignancies undergoing myeloablative chemotherapy followed by bone
	marrow transplant.
	3. Mobilization of progenitor cells into the peripheral blood stream for leukapheresis collection to be used after
Use Granulocyte Colony	myeloablative chemotherapy.
Stimulating Factor PA form	4. Treatment of congenital, cyclic, or idiopathic neutropenia in symptomatic patients.
	4. Treatment of congenitar, cyche, of idiopatine neutropenia in Symptomatic patients.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

	Updated 7/30/2012
Growth Hormone	Prior authorization is required for therapy with growth hormones. Payment for non-preferred growth hormones will be authorized only for cases in which there is documentation of previous trial and therapy failure with a preferred agent. All of the following criteria must be met for approval for prescribing of growth hormones: 1. Standard deviation of 2.0 or more below mean height for chronological age. 2. No intracranial lesion or tumor diagnosed by MRI. 3. Growth rate below five centimeters per year. 4. Failure of any two stimuli tests to raise the serum growth hormone level above ten nanograms per milliliter. 5. Annual bone age testing is required for the diagnosis of Growth Hormone Deficiency. A Bone age 14 to 15 years or less in females and 15 to 16 years or less in males is required. 6. Epiphyses open. Prior authorization will be granted for 12-month periods per patient as needed.
Use Growth Hormone PA	The following FDA approved indications for Growth Hormone therapy are considered not medically necessary and requests will be denied: Idiopathic Short Stature (ISS). If the request is for Zorbtive ® [somatropin (rDNA origin) for injection] approval will be granted for the treatment of Short Bowel Syndrome in patients receiving specialized nutritional support. Zorbtive® therapy should be used in conjunction
y .	with optimal management of Short Bowel Syndrome.
Hepatitis C Protease Inhibitors	Prior authorization is required for all oral hepatitis C protease inhibitors. Payment will be considered under the following conditions: 1. A diagnosis of hepatitis C genotype 1, and 2. Patient is 18 years of age or older, and 3. Administered in combination with peginterferon alfa and ribavirin. 4. HCV-RNA results are required at treatment week 4 for telaprevir (Incivek [™]). Additional prior authorization will be considered with documentation of response to treatment, measured by HCV-RNA levels. A maximum 12 weeks of therapy will be allowed for telaprevir (Incivek). 5. HCV-RNA results are required at treatment week 8, 12, and 24 (including lead in period) for boceprevir (Victrelis) and patient must not be a prior null responder to standart treatment. Additional prior authorizations will be considered with documentation of response to treatment, measured by HCV-RNA levels. Prior authorizations will be approved for a maximum of 24, 32, or 40 weeks of therapy with boceprevir (Victrelis) based on response.
Immunomodulators-	Prior authorization is required for topical immunomodulators. Payment for pimecrolimus (Elidel®) or tacrolimus
Topical	(Protopic®) 0.03% will be considered for non-immunocompromised patients two years of age and older and tacrolimus (Protopic®) 0.1% for patients 16 years of age and older when there is an adequate trial and therapy failure with two
Elidel [®] Protopic [®]	preferred topical corticosteroids. If criteria for coverage are met, requests will be approved for one tube per 90 days to ensure appropriate short-term and intermittent utilization of the medication. Quantities will be limited to 30 grams for use
Use Immunomodulators- Topical PA form	on the face, neck, and groin, and 60 grams or 100 grams for all other areas. The required trials may be overridden when documented evidence is provided that use of these agents would be medically contraindicated.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

	Updated 7/50/2012
Incretin Mimetic	Prior authorization is required for incretin mimetics (Byetta® & Victoza®). Payment will be considered under the
	following conditions: 1) Diagnosis of Type 2 diabetes mellitus, 2) Unless otherwise contraindicated, the member has not
Byetta [®]	achieved HbgA1C goals using a combination of two or more antidiabetic medications (metformin, sulfonylurea, or
Victoza [®]	thiazolidinedione) at maximum therapeutic doses. Initial authorizations will be approved for six months; additional prior
	authorizations will be considered on an individual basis after review of medical necessity and documented improvement in
Use Incretin Mimetic form	HbgA1C since the beginning of the initial prior authorization period.
Insulin, Pre-Filled	Prior authorization is required for pre-filled insulin pens. Prior authorization is granted when documentation indicates:
Pens	• The patient's visual or motor skills are impaired to such that they cannot accurately draw up their own insulin, and
	There is no caregiver available to provide assistance.
Use Pre-filled Insulin Pen	Prior authorization for non-preferred insulin pens will be authorized only for cases in which there is documentation of
PA form	previous trial and therapy failure with a preferred agent.
Isotretinoin (Oral)	Prior authorization is required for oral isotretinoin therapy. Payment will be approved for preferred oral isotretinoin
, ,	products for acne under the following conditions:
	1. There are documented trials and therapy failures of systemic antibiotic therapy and topical tretinoin therapy. Documented
	trials and therapy failures of systemic antibiotic therapy and topical tretinoin therapy are not required for approval for
	treatment of acne conglobata.
	2. Patients and providers must be registered in, and meet all requirements of, the iPLEDGE (www.ipledgeprogram.com)
	risk management program.
	nisk management program.
	Payment for non-preferred oral isotretinoin products will be authorized only for cases in which there is documentation of
Use Oral Isotretinoin PA form	trial(s) and therapy failure with a preferred agent(s). Initial authorization will be granted for up to 20 weeks. A minimum of
3	two months without therapy is required to consider subsequent
Ivacaftor _{TM}	Prior authorization is required for Kalydeco [™] (ivacaftor). Payment will be considered for patients when the following
(Kalydeco TM)	criteria are met:
	1. Patient is 6 years of age or older; and
	2. Has a diagnosis of cystic fibrosis with a G551D mutation in the CFTR gene as detected by a FDA-cleared CF mutation
	test; and
TA TA TM DAG	3. Prescriber is a CF specialist or pulmonologist; and
Use Kalydeco [™] PA form	4. Patient does not have one of the following infections: Burkholderia cenocepacia, dolosa, or Mycobacterium abcessus.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

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Ketorolac	Prior authorization is required for ketorolac tromethamine, a nonsteroidal anti-inflammatory drug indicated for short term
	(up to five days) management of moderately severe, acute pain. It is NOT indicated for minor or chronic conditions.
	This product carries a Black Box Warning . Initiate therapy with IV/IM and use oral ketorolac tromethamine only as a
	continuation therapy to ketorolac tromethamine IV/IM. The combined duration of use of IV/IM and oral is not to exceed
	five (5) days. Payment will be considered under the following conditions:
	1. For oral therapy, documentation of recent IM/IV ketorolac tromethamine injection including administration date and time, and the total number of injections given.
	2. Request falls within the manufacturer's dosing guidelines. Maximum oral dose is 40mg/day. Maximum IV/IM
	dose is 120mg/day. Maximum intranasal dose is 126mg/day. Maximum combined duration of therapy is 5 days per
	month.
	3. Diagnosis indicating moderately severe, acute pain.
	Requests for IV/IM and intranasal ketorolac must document previous trials and therapy failures with at least two preferred non-steroidal anti-inflammatory drugs at therapeutic doses.
Use Ketorolac PA form	
Lidocaine Patch	Prior authorization is required for topical lidocaine patches (Lidoderm®). Payment will be considered for a diagnosis of pain
(Lidoderm®)	associated with post-herpetic neuralgia following a previous treatment failure with a preferred agent at therapeutic dose
	from two of the following: tricyclic antidepressant, opioid, gabapentin, carbamazepine, or valproic acid. A maximum of 30
Use Lidocaine Patch	patches may be dispensed with the initial prescription to determine efficacy.
(Lidoderm®) PA form	
Linezolid (Zyvox®)	Prior authorization is required for Zyvox [®] . Payment for Zyvox [®] will be authorized when there is documentation that: 1. Prescriber is an infectious disease (ID) physician or has consulted ID physician (Telephone consultation is
	acceptable).
	2. Patient has an active infection and meets one of the following diagnostic criteria:
	 Vancomycin-resistant Enterococcus (VRE) and no alternatice regimens with documented efficacy are available and VRE is not in lower urinary tract**.
	 Methicillin-resistant Staph aureus (MRSA) and patient is intolerant to vancomycin*
	 Methicillin-resistant Staph epidermis (MRSE) and patient is intolerant to vancomycin*
	*Severe intolerance to vancomycin is defined as:
	 Severe rash, immune-complex mediated, determined to be directly related to vancomycin administration
	- Red-man's syndrome (histamine-mediated), refractory to traditional counter measures (e.g., prolonged IV infusion,
	premedicated with diphenhydramine)
	**VRE in lower urinary tract, considered to be pathogenic, may be treated with linezolid if severe renal insufficiency exists
Use Zyvox® PA form	and/or patient is receiving hemodialysis or has known hypersensitivity to nitrofurantoin.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

	Updated 7/50/2012
Milnacipran (Savella [™]) Use Chronic Pain Syndromes PA form	See Chronic Pain Syndromes Prior Authorization Criteria.
Modified Formulations	Payment for a non-preferred isomer, prodrug, or metabolite will be considered when the following criteria are met: 1. Previous trial with a preferred parent drug of the same chemical entity at a therapeutic dose that resulted in a partial response with a documented intolerance and
	2. Previous trial and therapy failure at a therapeutic dose with a preferred drug of a different chemical entity indicated to treat the submitted diagnosis if available. The required trials may be overridden when documented evidence is provided that use of these preferred agent(s) would be medically contraindicated.
Use Modified Formulations PA form	Prior authorization is required for the following modified dosage forms: Abilify Discmelt, Aplenzin, Aricept ODT, FazaClo, Invega, Metozolv ODT, Pristiq, Risperdal M-Tab, Suboxone Film, Trilipix, Xopenex, Zyprexa Zydis.
Muscle Relaxants Use Muscle Relaxant PA	Prior authorization is required for non-preferred muscle relaxants. Payment for non-preferred muscle relaxants will be authorized only for cases in which there is documentation of previous trials and therapy failures with at least three preferred muscle relaxants. Requests for carisoprodol will be approved for a maximum of 120 tablets per 180 days at a maximum dose of 4 tablets per day when the criteria for coverage are met. * If a non-preferred long-acting medication is requested, one trial must include the preferred immediate release product of the same chemical entity at a therapeutic dose, unless
form	evidence is provided that use of these products would be medically contraindicated.
Narcotic Agonist- Antagonist Nasal Sprays	Prior authorization is required for narcotic agonist-antagonist nasal sprays. For consideration, the diagnosis must be supplied. If the use is for the treatment of migraine headaches, documentation of current prophylactic therapy or documentation of previous trials and therapy failures with two different prophylactic medications must be provided. There must also be documented treatment failure or contraindication to triptans for the acute treatment of migraines. For other pain conditions, there must be documentation of treatment failure or contraindication to oral administration.
	Payment for non-preferred narcotic agonist-antagonist nasal sprays will be authorized only for cases in which there is documentation of previous trial and therapy failure with a preferred agent.
Use Narcotic Agonist/Antagonist Nasal Spray PA form	Quantities are limited to 2 bottles or 5 milliliters per 30 days. Payment for narcotic agonist-antagonist nasal sprays beyond this limit will be considered on an individual basis after review of submitted documentation.
Nebivolol (Bystolic®) Use Nebivolol (Bystolic®) PA form	Prior authorization is required for Bystolic [®] . Payment will be considered in cases where there are documented trials and therapy failures with two preferred cardio-selective beta-blockers of a different chemical entity at a therapeutic dose. The required trials may be overridden when documented evidence is provided that the use of these agents would be medically contraindicated.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

	Updated 7/50/2012
Nicotine	Prior Authorization is required for over-the-counter nicotine replacement patches, gum, or lozenges, and prescription
Replacement	nicotine nasal spray or inhaler. Requests for authorization must include:
Therapy	1) Diagnosis of nicotine dependence and referral to the Quitline Iowa program for counseling.
	2) Confirmation of enrollment in the Quitline Iowa counseling program is required for approval. Continuation
	therapy is available only with documentation of ongoing participation in the Quitline Iowa program.
	3) Approvals will only be granted for patients eighteen years of age and older.
	4) The maximum allowed duration of therapy is twelve weeks total combined therapy within a twelve-month
	period.
	5) Patients may receive nicotine replacement patches in combination with one of the oral nicotine replacement
	products (gum or lozenges). A maximum quantity of 14 nicotine replacement patches and 110 pieces of
	nicotine gum or 144 nicotine lozenges may be dispensed with the initial prescription. Subsequent prescription
	refills will be allowed to be dispensed as a 4 week supply at one unit per day of nicotine replacement patches
	and 330 pieces of nicotine gum or 288 nicotine lozenges.
	6) Requests for non-preferred nicotine replacement products will be considered after documentation of previous
	trials and intolerance with a preferred oral and preferred topical nicotine replacement product. A maximum
Use Nicotine	quantity of 168 nicotine inhalers or 40ml nicotine nasal spray may be dispensed with the initial prescription.
Replacement Therapy PA	Subsequent prescription refills will be allowed to be dispensed as a 4 week supply at 336 nicotine inhalers or
form	80ml of nicotine nasal spray.
	7) The 72-hour emergency supply rule does not apply for drugs used for the treatment of smoking cessation.
Non-Parenteral	Prior authorization is required for non-parenteral vasopressin derivatives of posterior pituitary hormone products. Payment
Vasopressin	for preferred non-parenteral vasopressin derivatives of posterior pituitary hormone products will be authorized for the
Derivatives of	following diagnoses:
Posterior Pituitary	1. Diabetes Insipidus.
Hormone Products	2. Hemophilia A.
	3. Von Willebrand's disease.
	Payment for oral vasopressin derivatives of posterior pituitary hormone products used in the treatment of primary nocturnal
Use Non-Parenteral	enuresis will be authorized for patients who are six years of age or older for periods of six months. Approvals will be
Vasopressin Deriv. of	granted for subsequent six-month periods only after a drug-free interval to assess the need for continued therapy. Payment
Posterior Pituitary	for non-preferred non-parenteral vasopressin derivatives will be authorized only for cases in which there is documentation
Hormone Products PA form	of trial and therapy failure with the preferred agent.
Non-Preferred Drug	Prior authorization is required for non-preferred drugs as specified on the Iowa Medicaid Preferred Drug List. Payment for
II N D C ID	a non-preferred medication will be authorized only for cases in which there is documentation of previous trial and therapy
Use Non-Preferred Drug PA form	failure with the preferred agent, unless evidence is provided that use of these agents would be medically contraindicated.
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The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

	Cptatet 1/30/2012
Nonsteroidal Anti-	Prior authorization is required for all non-preferred nonsteroidal anti-inflammatory drugs (nsaids) and COX-2 inhibitors.
inflammatory Drugs	Prior authorization is not required for preferred nonsteroidal anti-inflammatory drugs or COX-2 inhbitors.
	 Requests for a non-preferred nsaid must document previous trials and therapy failures with at least three preferred nsaids.
	 Requests for a non-preferred COX-2 inhibitor must document previous trials and therapy failures with three preferred nsaids, two of which must be a preferred COX-2 preferentially selective nsaid.
	3. Requests for a non-preferred topical nsaid must document previous trials and therapy failures with three preferred nsaids. The trials must include two preferred COX-2 preferentially selective nsaids and the oral drug of the same chemical entity. In addition, the use of a topical delivery system must be deemed medically necessary.
T N G HAA	4. Requests for a non-preferred extended release nsaid must document previous trials and therapy failures with three preferred nsaids, one of which must be the preferred immediate release nsaid of the same chemical entity at a therapeutic dose that resulted in a partial response with a documented intolerance.
Use Non-Steroidal Anti- inflammatory Drug PA form	The required trials may be overridden when documented evidence is provided that use of these agents would be medically contraindicated.
Omalizumab	Prior authorization is required for Xolair [®] . Payment for Xolair [®] will be authorized for patients 12 and older when there is a
(Xolair®)	diagnosis of moderate to severe persistent asthma and documentation of previous trial and therapy failure with therapeutic doses of inhaled steroids.
Use Xolair® PA form	Extended release oxycodone/OxyContin [®] is non-preferred except for patients being treated for cancer related pain.Prior
Oxycodone CR/ER (OxyContin [®])	authorization at any dose twice daily for cancer related pain will be approved. For all other diagnoses, payment will be considered under the following conditions:
	1. There is documentation of previous trials and therapy failures with two (2) chemically distinct preferred long-acting narcotics (such as extended-release morphine sulfate and methadone) at therapeutic doses, and
Use OxyContin® PA form	2. A trial and therapy failure with fentanyl patch at maximum tolerated doses, and
	3. A signed chronic opioid therapy management plan between the prescriber and patient must be included with the prior authorization, and
	4. The prescriber must review the patient's use of controlled substances on the Iowa Prescription Monitoring Program website at https://pmp.iowa.gov/IAPMPWebCenter/ prior to requesting prior authorization.
	5. Requests will only be considered for 12 hour dosing.
	The required trials may be overridden when documented evidence is provided that use of these agents would be medically contraindicated.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

	Updated 7/30/2012
Palivizumab (Synagis [®])	Prior authorization is required for therapy with palivizumab. Prior authorizations will be approved for a maximum of five doses per patient. No allowances will be made for a sixth dose. Some patients may receive a maximum of three doses, dependent on gestational and chronological age at the start of the RSV season. Payment for palivizumab will be considered for patients who meet one of the following criteria: Chronic Lung Disease (CLD)
	 Patient is less than 24 months of age at start of therapy and has chronic lung disease of prematurity (i.e. bronchopulmonary dysplasia) requiring medication (bronchodilator, corticosteroid, or diuretic therapy) or oxygen within six months before the anticipated start of RSV season. Prematurity
	 Patient is less than 12 months of age at start of therapy with a gestational age of less than 29 weeks. Patient is less than 6 months of age at start of therapy with a gestational age of 29 weeks through 31 weeks. Patient is less than 3 months of age at start of therapy or born during the RSV season with a gestational age of 32 weeks through 34 weeks and has one of two risk factors. Risk factors include: day care attendance or siblings less than 5 years of age in household. Doses will be limited to a maximum of 3 doses or until patient reaches 90 days of age, which ever comes first.
Use Palivizumab PA form	 Severe Neuromuscular Disease or Congenital Abnormalities Patient is 12 months of age or younger at the start of therapy and has either severe neuromuscular disease or congenital abnormalities of the airway that compromises handling of respiratory secretions. Congenital Heart Disease (CHD) Patient is less than 24 months of age at start of therapy and has hemodynamically significant congenital heart disease further defined by any of the following: Receiving medication to control congestive heart failure, moderate to severe pulmonary hypertension, or cyanotic congenital heart disease. Severe Immunodeficiency Patient is less than 24 months of age at start of therapy and has severe immunodeficiencies (e.g., severe combined)
Pregabalin (Lyrica®) Use Chronic Pain Syndromes PA form	immunodeficiency or advanced acquired immunodeficiency syndrome). See Chronic Pain Syndromes Prior Authorization Criteria.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

	Cptatea 7/30/2012
Proton Pump Inhibitors	Prior authorization is not required for the preferred proton pump inhibitors (PPI) for a cumulative 60-days of therapy per 12-month period. Prior authorization will be required for all non-preferred proton pump inhibitors as indicated on the Iowa Medicaid Preferred Drug List beginning the first day of therapy. Payment for a non-preferred proton pump inhibitor will be authorized only for cases in which there is documentation of previous trials and therapy failures with three preferred products. Prior authorization is required for any PPI usage longer than 60 days or more frequently than one 60-day course per 12-month period. The 12-month period is patient specific and begins 12 months before the requested date of prior authorization. Payment for usage beyond these limits will be authorized for cases in which there is a diagnosis of:
	 Specific Hypersecretory conditions (Zollinger-Ellison syndrome, systemic mastocytosis, multiple endocrine adenomas). Barrett's esophagus. Erosive esophagitis Symptomatic gastroesophageal reflux after documentation of previous trials and therapy failure with at least one
	histamine H2-receptor antagonist at full therapeutic doses. Requests for PPIs exceeding one unit per day will be considered after documentation of a therapeutic trial and therapy failure with concomitant use of once daily PPI dosing and a bedtime dose of a histamine H2-receptor antagonist. Upon failure of the combination therapy, subsequent requests for PPIs exceeding one unit per day will be considered on a short term basis (up to 3 months). After the three month period, a retrial of the recommended once daily dosing will be required on an annual basis for those patients continuing to need doses beyond one unit per day.
Use Proton Pump Inhibitor PA form	5. Recurrent peptic ulcer disease after documentation of previous trials and therapy failure with at least one histamine H2-receptor antagonist at full therapeutic doses and with documentation of either failure of Helicobacter pylori treatment or a negative Helicobacter pylori test result.
Pulmonary Arterial Hypertension Agents Use Pulmonary Arterial Hypertension Agents PA form	Prior Authorization is required for agents used to treat pulmonary hypertension. Payment will be approved under the following conditions: 1. Diagnosis of pulmonary arterial hypertension
Quantity Limit Override Use Quantity Limit Override PA form	Designated drugs are limited to specific quantity limitations. These drugs are identified on the Iowa Medicaid Quantity Limit Chart posted on the website www.iowamedicaidpdl.com under the Billing/Quantity Limits tab. Providers should submit a Prior Authorization request for override consideration.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

	TM Cpdatcd 1/30/2012
Roflumilast	Prior authorization is required for roflumilast (Daliresp [™]). Payment will be considered for patients 18 years of age or older
(Daliresp [™])	when the following is met:
	1. A diagnosis of severe COPD with chronic bronchitis as documented by spirometry results, and
	2. A smoking history of ≥ 20 pack-years, and
	3. Currently on a long-acting bronchodilator in combination with an inhaled corticosteroid with documentation of
	inadequate control of symptoms, and
Use Roflumilast	4. A history of at least one exacerbation in the past year requiring treatment with oral glucocorticosteroids.
(Daliresp [™]) PA form	The required trials may be overridden when documented evidence is provided that the use of these agents would be
	medically contraindicated.
Sedative/Hypnotics-	Preferred agents are available without prior authorization (PA). Although intermittent therapy is recommended, quantity
Non-Benzodiazepine	limits will allow for 30 tablets per 30 days supply without PA for preferred medications.
Tion Benzouluzepine	Prior authorization is required for all non-preferred non-benzodiazepine sedative/hypnotics. Payment for non-preferred non-
	benzodiazepine sedative/hypnotics will be authorized only for cases in which there is documentation of a previous trial and
	therapy failure with the preferred agent(s). Payment for non-preferred non-benzodiazepine sedative/hypnotics will be
	considered when there is:
	1) A diagnosis of insomnia,
	2) Medications with a side effect of insomnia (i.e. stimulants) are decreased in dose, changed to a short acting
	product, and/or discontinued,
	3) Enforcement of good sleep hygiene is documented.
Use Sedative/Hypnotics-	4) All medical, neurological, and psychiatric disease states causing chronic insomnia are being adequately treated
Non-Benzodiazepine PA	with appropriate medication at therapeutic doses.
form	5) Patient has a documented trial and therapy failure with zaleplon.
Selected Brand Name	Prior authorization is required for selected brand-name drugs, as determined by the Department, for which there is available
Drugs	an "A" rated bioequivalent generic product as determined by the Federal Food and Drug Administration, unless the brand
Drugs	drug has been designated by the Department as preferred (payable) under the Iowa Medicaid Preferred Drug List (PDL).
	For prior authorization to be considered, the prescriber must submit a completed Selected Brand Name PA form and Iowa
	Medicaid MedWatch form with:
	1. Documentation of trials and therapy failures with two different generic manufacturers of the same chemical entity. If an
	allergy to an inactive component is suspected, the second trial must be with a generic product that does not contain the
	allergen, if available.
	2. Documentation of the failure must include the specific adverse reaction as defined by the FDA (See Section B of the
	MedWatch form). Intolerances, such as nausea or vomiting, to the generic drug will not be considered as a basis for
Use Selected Brand Name	approval.
PA forms	Trials may be overridden when evidence is provided that use of the generic product would be medically contraindicated.

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

Serotonin 5-HT1-	Prior authorization is required for preferred serotonin 5-HT1-receptor agonists for quantities exceeding 12 unit doses of
receptor Agonists	tablets, syringes or sprays per 30 days. Payment for serotonin 5-HT1-receptor agonists beyond this limit will be considered
	on an individual basis after review of submitted documentation. Prior authorization will be required for all non-preferred
	serotonin 5-HT1-receptor agonists as indicated on the Iowa Medicaid Preferred Drug List beginning the first day of therapy.
	Payment for non-preferred serotonin 5-HT1-receptor agonists will be authorized only for cases in which there is
	documentation of previous trials and therapy failures with two preferred agents. Requests for non-preferred combination
	products may only be considered after documented separate trials and therapy failures with the individual ingredients. For
	consideration, the following information must be supplied:
	1. The diagnosis requiring therapy.
Use Serotonin 5-HT1-	2. Documentation of current prophylactic therapy or documentation of previous trials and therapy failures with two
receptor Agonists PA form	different prophylactic medications.
Short Acting	Prior authorization is required for all non-preferred short acting narcotics. Payment will be considered for cases in which
Narcotics	there is documentation of previous trials and therapy failures with three (3) chemically distinct preferred short acting
	narcotics (based on narcotic ingredient only) at therapeutic doses, unless evidence is provided that use of these products
Use Short Acting Narcotics	would be medically contraindicated.
PA form	
Smoking Cessation	Prior Authorization is required for varenicline (Chantix®) or bupropion SR that is FDA approved for smoking cessation.
Therapy-Oral	Requests for authorization must include:
	1) Diagnosis of nicotine dependence and referral to the Quitline Iowa program for counseling.
<i>Chantix</i> ®	2) Confirmation of enrollment and ongoing participation in the Quitline Iowa counseling program is required for
Bupropion SR	approval and continued coverage.
	3) Approvals will only be granted for patients eighteen years of age and older.
	4) The duration of therapy is initially limited to twelve weeks within a twelve-month period. For patients who
	have successfully stopped smoking at the end of 12 weeks, an additional course of 12 weeks treatment will be
	considered with a prior authorization request. The maximum duration of approvable therapy is 24 weeks within
	a twelve-month period.
	5) Requests for varenicline to be used in combination with bupropion SR that is FDA indicated for smoking
	cessation or nicotine replacement therapy will not be approved.
Use Smoking Cessation	6) The 72-hour emergency supply rule does not apply for drugs used for the treatment of smoking cessation
Therapy-Oral PA form	e, and the second control of the second seco

The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Updated 7/30/2012

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Sodium Oxybate	Prior authorization is required for sodium oxybate (Xyrem [®]). Payment will be considered for patients 16 years of age or
(Xyrem [®])	older under the following conditions:
	1.A diagnosis of cataplexy associated with narcolepsy verified by a recent sleep study (including PSG, MSLT, and
	ESS) and previous trial and therapy failure with one of the following tricyclic antidepressants: clomipramine,
	imipramine, or protriptyline.
	2.A diagnosis of excessive daytime sleepiness associated with narcolepsy verified by a recent sleep study (including
	PSG, MSLT, and ESS) and previous trials and therapy failures at a therapeutic dose with a preferred amphetamine
	and non-amphetamine stimulant.
	3. Requests for patients with a prior history of substance abuse, concurrent use with a sedative hypnotic, or a
	semialdehyde dehydrogenase deficiency will not be considered.
<u>Use Sodium Oxybate</u>	The required trials may be overridden when documented evidence is provided that the use of these agents would be
(Xyrem [®]) PA form	medically contraindicated.
Thrombopoietin	Payment for a preferred thrombopoietin receptor agonist will only be considered for cases in which there is a diagnosis of
Receptor Agonists	chronic immune thrombocytopenic purpura (ITP) including documentation of an insufficient response to a corticosteroid,
	an immunoglobulin, or the patient has undergone a splenectomy. Payment for a non-preferred thrombopoietin receptor
Use Thrombopoietin	agonist will be considered following documentation of a recent trial and therapy failure with a preferred thrombopoietin
Receptor Agonists PA form	receptor agonist unless such a trial would be medically contraindicated.
Topical Retinoids for	Prior authorization is required for all prescription topical retinoid products. Payment for prescription topical retinoid
Acne	products will be considered under the following conditions:
	1. Previous trial and therapy failure with a preferred over-the-counter benzoyl peroxide product, and
	2. Previous trials and therapy failures with two preferred topical and/or oral antibiotics for the treatment of mild to
	moderate acne (non-inflammatory and inflammatory), and drug-induced acne.
	3. Payment for non-preferred topical retinoid products will be authorized only for cases in which there is
	documentation of previous trial and therapy failure with a preferred agent.
	4. Trials and therapy failure will not be required for those patients presenting with a preponderance of comedonal
	acne.
	5. Skin cancer, lamellar ichthyosis, and Darier's disease diagnoses will receive automatic approval for lifetime use
	of topical retinoid products.
	6. Requests for non-preferred combination products may only be considered after documentation of separate trials
	and therapy failures with the individual ingredients.
	7. Requests for tazorac for a psoriasis diagnosis may only be considered after documentation of a previous trial and
Use Topical Retinoids for	therapy failure with a preferred topical antipsoriatic agent.
Acne PA form	The required trials may be overridden when documented evidence is provided that the use of these agents would be
-	medically contraindicated.
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The drug prior authorization unit will consider other conditions as listed in the compendia on an individual basis after reviewing documentation submitted regarding the medical necessity. All required trials must be of appropriate dose and duration for the indication and must be documented by the prescriber, on the request for prior authorization form, including dates, dose, and nature of failure.

Vilazodone	Prior authorization is required for Viibryd [™] . Requests for doses above the manufacturer recommended dose will not be
(Viibryd [™])	considered. Payment will be considered for patients when the following criteria are met:
	1. The patient has a diagnosis of Major Depressive Disorder (MDD) and is 18 years of age or older; and
	2. Documentation of a previous trial and therapy failure at a therapeutic dose with one preferred generic SSRI; and
	3. Documentation of a previous trial and therapy failure at a therapeutic dose with one preferred generic SNRI; and
	4. Documentation of a previous trial and therapy failure at a therapeutic dose with an additional generic
Use Vilazodone (Viibryd TM)	antidepressant from any class.
PA form	The required trials may be overridden when documented evidence is provided that the use of these agents would be
	medically contraindicated.
Vitamins, Minerals	Payment for vitamins, minerals and multiple vitamins for treatment of specific conditions will be approved when there is a
and Multiple	diagnosis of specific vitamin or mineral deficiency disease or for patients under 21 years of age if there is a diagnosed
Vitamins	disease which inhibits the nutrition absorption process as a secondary effect of the disease. (Prior approval is not required
	for prescribed multi-vitamins with or without iron or vitamin D supplements for patients under 12 months of age or a
Use Vitamin/Mineral PA	prescription product primarily classified as a blood modifier, if that product does not contain more than three
form	vitamins/minerals or for products principally marketed as prenatal vitamin-mineral supplements.)
Vusion [™] Ointment	Prior Authorization is required for Vusion [™] Ointment. Payment will only be considered for cases in which there is
	documentation of previous trials and therapy failures with 1) over-the-counter miconazole 2% cream (payable with a
II II TM O' (DA	prescription) AND 2) nystatin cream or ointment, unless evidence is provided that use of these agents would be medically
Use Vusion [™] Ointment PA form	contraindicated.