





FAX Completed Form To 1.833.404.2392

Prescriber Help Desk 1.833.587.2012

Online

covermymeds.com/main/prior-authorization-forms/

Request for Prior Authorization Gonadotropin-Releasing Hormone (GnRH) Receptor Antagonist, Oral

(PLEASE PRINT - ACCURACY IS IMPORTANT)

IA Medicaid Member ID #	Patient name	DOB			
Patient address					
Provider NPI	Prescriber name	Phone			
Prescriber address		Fax			
Pharmacy name	Address	Phone			
·					
Prescriber must complete all information above. It must be legible, correct, and complete or form will be returned.					
Pharmacy NPI	Pharmacy fax	NDC			

Prior authorization is required for oral gonadotropin-releasing hormone (GnRH) antagonists. Payment for non-preferred oral GnRH antagonists may be considered only for cases in which there is documentation of a previous trial and therapy failure with the preferred agent. Payment will be considered for patients when the following is met:

- 1) Pregnancy has been ruled out; and
- 2) Patient does not have osteoporosis; and
- 3) Request adheres to all FDA approved labeling for requested drug, including age, dosing, contraindications, warnings and precautions, drug interactions, and use in specific populations; and
- 4) Requests for elagolix (Orilissa) will be considered under the following conditions:
 - a. Patient has a diagnosis of moderate to severe pain associated with endometriosis; and
 - b. Patient has documentation of a previous trial and therapy failure with at least one preferred oral NSAID and at least one preferred 3-month course of a continuous hormonal contraceptive taken concurrently; and
 - c. Patient has documentation of a previous trial and therapy failure with a preferred GnRH agonist.
 - d. Initial requests will be considered for 3 months. Additional requests will be considered upon documentation of improvement of symptoms.
 - e. Requests will be considered for a maximum of 24 months for the 150mg dose and 6 months for the 200mg dose; or
- 5) Requests for elagolix, estradiol, and norethindrone acetate; elagolix (Oriahnn) or relugolix, estradiol, norethindrone acetate (Myfembree) will be considered under the following conditions:
 - a. Patient is premenopausal; and
 - b. Patient has a diagnosis of heavy menstrual bleeding associated with uterine leiomyomas (fibroids); and
 - c. Patient has documentation of a previous trial and therapy failure with at least one preferred 3-month course of a continuous hormonal contraceptive; and
 - d. Patient has documentation of a previous trial and therapy failure with tranexamic acid.
 - e. Initial requests will be considered for 6 months. Additional requests will be considered upon documentation of improvement in symptoms.
 - f. Requests will be considered for a maximum of 24 months of treatment.

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Preferred	Non-Preferred					
☐ Oriahnn		Orilissa				
Strength	Dosage Instruc	ctions	Quantity	Days Supply		
Diagnosis:						
☐ Initial Requests:						
Has pregnancy been ru	led out? Yes	☐ No Date	of pregnancy test:_			
Does patient have oste	oporosis?	☐ No				
Does patient have seve	ere hepatic impairment?	Yes N	0			
Is patient taking a strong organic anion transporting polypeptide (OATP) 1B1 inhibitor (e.g., cyclosporine and gemfibrozil)?						
Treatment Failures:						
Preferred Oral NSAID Tria	al:					
Name/dose:			Trial dates:			
Failure reason/medical con	traindication:					
Preferred Continuous Ho Name/dose: Failure reason/medical con						
Preferred GnRH Agonist	Trial:					
Name/dose:			_ Trial dates:			
Failure reason/medical con	traindication:					

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Oriahnn & Myfembree			
Is patient premenopausal?	☐ Yes ☐ No		
Treatment Failures:			
Preferred Continuous Hormonal C	Contraceptive Trial:		
Name/dose:		Trial dates:	
Failure reason/medical contraindicat			
Tranexamic Acid Trial:			
Name/dose:		Trial dates:	
Failure reason/medical contraindicat			
Reason for use of Non-Preferred of	drug requiring prior app	proval:	
Renewal Requests:			
Provide documentation of improvem	ent in symptoms:		
Treatment start date:			
Attach lab results and other docu	mentation as necessary	<i>/</i> .	
Prescriber signature (Must match prescri	iber listed above.)	Date of submission	

IMPORTANT NOTE: In evaluating requests for prior authorization the consultant will consider the treatment from the standpoint of medical necessity only. If approval of this request is granted, this does not indicate that the member continues to be eligible for Medicaid. It is the responsibility of the provider who initiates the request for prior authorization to establish by inspection of the member's Medicaid eligibility card and, if necessary by contact with the county Department of Human Services, that the member continues to be eligible for Medicaid.

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