UTERINE DISORDERS PRIOR AUTHORIZATION REQUEST FORM



OptumRx
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(magaille						Community Plan
]				
Note: This form must	be complete	ed by the p	rescribin	g pro	ovider.	
*:	*All sections	must be co	ompleted	or t	he request will be returned**	
Patient's Medicaid #				Date of Birth / / /		
Patient's Name				Prescriber's Name		
Prescriber's IN License #				Specialty		
Prescriber's NPI #				Prescriber's Signature		
Return Fax				Return Phone #		
Check box if requesting retro-active PA				Date(s) of service requested for retro-active eligibility (if applicable):		
Note: Submit PA requests for retroactive claims (dates of service prior to eligibility determination, but within established eligibility timelines) with dates of service prior to 30 calendar days of submission separately from current PA requests (dates of service 30 calendar days or less and going forward).						
Requested Medic	ation	Strength	Quanti	ity	Dosage Regi	men
PA requirements fo	or MYFEM	BREE (re	luaolix/	estr	adiol/norethindrone ace	etate):
1. Member is 18 years o						
	associated v	vith uterine le	•	•	oroids) in premenopausal fema sis in premenopausal females	
3. Negative pregnancy test in the past 30 days* \square Yes \square No						
4. Laboratory tests confi	rming no hep	oatic disease	e in the pa	st 30) days* ☐ Yes ☐ No	
 5. Provider attests that member has none of the following contraindications to therapy: Yes No Current diagnosis of, risk factors for, or previous history of thromboembolic disorders or vascular events Current diagnosis or history of breast cancer or other hormone-sensitive malignancies OR increased risk factors for hormone-sensitive malignancies Diagnosis of osteoporosis Undiagnosed abnormal uterine bleeding 						
g		3				

If no , please specify contraindication and medical justification for use:
Prescriber Signature:
6. Requested dose is 1 tablet (40/1/0.5 mg) per day ☐ Yes ☐ No
If no , please explain
 7. Previous trial and failure of one of the following: Hormonal contraceptives/therapy (oral tablets, vaginal ring, patch, intrauterine contraception (IUD)) for menorrhagia associated with uterine leiomyomas indication ONLY Yes No Hormonal contraceptives/therapy (oral tablets, vaginal ring, patch, IUD) AND NSAID therapy for endometriosis indication ONLY Yes No
If no , please provide medical justification:
8. Member will not be exceeding 24 months of therapy per lifetime with Myfembree (relugolix/estradiol/norethindrone acetate) Yes No
If yes , provide medical justification for continued use beyond 24 months and date range or number of months member has received therapy thus far:
*Note: Chart documentation will need to be provided for questions indicated with asterisk
PA requirements for ORIAHNN (elagolix/estradiol/norethindrone acetate):
1. Member is 18 years of age or older ☐ Yes ☐ No
2. Diagnosis of menorrhagia associated with uterine leiomyomas (fibroids) in premenopausal females ☐ Yes ☐ No
3. Negative pregnancy test in the past 30 days* \square Yes \square No
4. Laboratory tests confirming no hepatic disease in the past 30 days* \square Yes \square No
 5. Provider attests that member has none of the following contraindications to therapy: Yes No Concurrent use of organic anion transporting polypeptide (OATP)1B1 inhibitors that are known or expected to significantly increase elagolix plasma concentrations (e.g., cyclosporine, gemfibrozil) Current diagnosis of, risk factors for, or previous history of thromboembolic disorders or vascular events Current diagnosis or history of breast capper or other hormone sensitive malignancies OR increased
 Current diagnosis or history of breast cancer or other hormone-sensitive malignancies OR increased risk factors for hormone-sensitive malignancies Diagnosis of osteoporosis
 Undiagnosed abnormal uterine bleeding If no, please specify contraindication and medical justification for use:

Prescrib	er Signature:
6. Reque	sted dose is 2 capsules (1 x 300/1/0.5 mg; 1 x 300 mg) per day \square Yes \square No
lf no ,	please explain
	us trial and failure of hormonal contraceptives/therapy (oral tablets, vaginal ring, patch, and terine contraception) $\ \square$ Yes $\ \square$ No
lf no , ple	ase provide medical justification:
	er will not be exceeding 24 months of therapy per lifetime with elagolix/estradiol/norethindrone acetate by ☐ Yes ☐ No
	ovide medical justification for continued use beyond 24 months and date range or number of months has received therapy thus far:
*Note: C	hart documentation will need to be provided for questions indicated with asterisk
PA req	
•	uirements for ORILISSA (elagolix):
	er is 18 years of age or older Yes No
1. Memb	
1. Memb 2. Select	er is 18 years of age or older Yes No one of the following diagnoses: Moderate to severe pain associated with endometriosis with co-existing endometriosis-related dyspareunia AND dose does not exceed 400 mg daily (6-month approval maximum) Moderate to severe pain associated with endometriosis AND requested dose does not exceed 150
1. Memb 2. Select 3. Negati 4. Labora	er is 18 years of age or older Yes No one of the following diagnoses: Moderate to severe pain associated with endometriosis with co-existing endometriosis-related dyspareunia AND dose does not exceed 400 mg daily (6-month approval maximum) Moderate to severe pain associated with endometriosis AND requested dose does not exceed 150 mg daily (1 year approval)
1. Memb 2. Select 3. Negati 4. Labora Fig. 5. Provid	er is 18 years of age or older
1. Memb 2. Select 3. Negati 4. Labora Final 5. Provid	er is 18 years of age or older \ Yes \ No one of the following diagnoses: \[\] Moderate to severe pain associated with endometriosis with co-existing endometriosis-related dyspareunia AND dose does not exceed 400 mg daily (6-month approval maximum) \[\] Moderate to severe pain associated with endometriosis AND requested dose does not exceed 150 mg daily (1 year approval) ve pregnancy test in the past 30 days* \ Yes \ No atory tests confirming no hepatic disease worse than Child-Pugh class B in the past 30 days* Please indicate Child-Pugh classification if applicable: \[\] Child-Pugh class A \ \ Child-Pugh class B \ N/A Note: members with Child-Pugh class B will be limited to 150 mg daily dose for a maximum of 6 months respective of indication er attests that member has none of the following contraindications to therapy: \ Yes \ No Diagnosis of osteoporosis Concurrent use of organic anion transporting polypeptide (OATP)1B1 inhibitors that are known or

6. Previous trial and failure of hormonal contraceptives/therapy (oral tablets, vaginal ring, patch, and intrauterine contraception) AND NSAID therapy ☐ Yes ☐ No
If no , please provide medical justification:
7. Member will not be exceeding 24 months of therapy per lifetime with elagolix \square Yes \square No
If yes , provide medical justification for continued use beyond 24 months and date range or number of month member has received therapy thus far:
*Note: Chart documentation will need to be provided for questions indicated with asterisk

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