SENTARA COMMUNITY PLAN (MEDICAID)

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid)</u> on this request. All other information may be filled in by office staff; <u>fax to 1-800-750-9692</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

The Sentara Health Plans Oncology Program is administered by OncoHealth

❖ For any oncology indications, the most efficient way to submit a prior authorization request is through the OncoHealth OneUM Provider Portal at https://oneum.oncohealth.us. Fax to 1-800-264-6128. OncoHealth can also be contacted by Phone: 1-888-916-2616.

<u>Drug Requested:</u> Gonadotropin-releasing Hormone Agonists (GnRH) (Pharmacy)

Preferred Drugs				
Camcevi [™] (leuprolide) 42 mg (6-month)		Eligard® (leuprolide) 22.5 mg (3-month)		Eligard® (leuprolide) 30 mg (4-month)
Eligard® (leuprolide) 45 mg (6-month)		Leuprolide Depot 22.5 mg (3-month) [vial]		Lupron Depot® (leuprolide acetate) 11.25 mg (3-month)
Lupron Depot® (leuprolide acetate) 22.5 mg (3-month)		Lupron Depot® (leuprolide acetate) 30 mg (4-month)		Lupron Depot® (leuprolide acetate) 45 mg (6-month)
Lupron Depot-Ped® (leuprolide acetate) 11.25 mg (3-month)		Lupron Depot-Ped® (leuprolide acetate) 30 mg (3- month)		Lupron Depot-Ped® (leuprolide acetate) 45 mg (6-month)
Lutrate Depot (leuprolide acetate) 22.5 mg (3- month) [vial]		Supprelin® LA (histrelin) 50 mg (12-month) *Requires authorization under medical benefit		Trelstar® (triptorelin pamoate) 11.25 mg (3-month)
Trelstar® (triptorelin pamoate) 22.5 mg (3-month)		Vantas [™] (histrelin) 50mg (12- month) *Requires authorization under medical benefit		

Non-Preferred Drugs				
Eligard® (leuprolide) 7.5 mg (1-month)		Fensolvi® (leuprolide) 45 mg (6-month)		Leuprolide acetate 5 mg/mL SubQ Solution
Lupron Depot® (leuprolide acetate) 3.75 mg (1-month)		Lupron Depot ® (leuprolide acetate) 7.5 mg (1-month)		Lupron Depot-Ped® (leuprolide acetate) 7.5 mg (1-month)
Lupron Depot-Ped® (leuprolide acetate) 11.25 mg (1- month)		Lupron Depot-Ped® (leuprolide acetate) 15 mg (1- month)		Synarel® (nafarelin) 2 mg/ml (dosing of 1600 mcg to 1800 mcg per day more than the 400 mcg to 800 mcg per day for endometriosis)
Trelstar® (triptorelin pamoate) 3.75 mg (1-month)		Triptodur® (triptorelin) 22.5 mg (6-month)		Zoladex® (goserelin) 3.6 mg (1-month) *Requires authorization under medical benefit
Zoladex® (goserelin) 10.8 mg (3-month) *Requires authorization under medical benefit				

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.				
Member Name:				
Member Sentara #:				
Prescriber Name:				
Prescriber Signature:				
Office Contact Name:				
Phone Number:	Fax Number:			
NPI #:				
DRUG INFORMATION: Authoriz				
Drug Name/Form/Strength:				
Dosing Schedule:	Length of Therapy:			
Diagnosis:	ICD Code, if applicable:			
Weight (if applicable):	Date weight obtained:			

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A. Length of Authorization

- Endometriosis: Coverage will be provided for 6 months and is eligible for conditional renewal
- Uterine leiomyomata (fibroids): Coverage will be provided for 6 months and is eligible for conditional renewal
- Gender Dysphoria: Approval and renewal authorizations will be provided for 12 months
- All other indications: Coverage will be provided for 12 months and may be renewed

B. Quantity Limits:

Drug Name	Strength	Quantity	Day Supply
Camcevi	42 mg	1 injection	180 days
Leuprolide acetate SubQ solution	5 mg/mL (1 mg/0.2 mL multi-dose vial)	2 vials	28 days
Leuprolide Depot 3-month	22.5 mg	1 vial	84 days
Lupron Depot 1-month	3.75 mg, 7.5 mg	1 injection	28 days
Lupron Depot 3-month	11.25 mg, 22.5 mg	1 injection	84 days
Lupron Depot 4-month	30 mg	1 injection	112 days
Lupron Depot 6-month	45 mg	1 injection	168 days
Lupron Depot-Ped 1-month	7.5 mg, 11.25 mg, 15 mg	1 injection	28 days
Lupron Depot-Ped 3-month	11.25 mg, 30 mg	1 injection	84 days
Lupron Depot-Ped 6-month	45 mg	1 injection	168 days
Lutrate Depot 3-month	22.5 mg	1 vial	84 days
Eligard 1-month	7.5 mg	1 injection	28 days
Eligard 3-month	22.5 mg	1 injection	84 days
Eligard 4-month	30 mg	1 injection	112 days
Eligard 6-month	45 mg	1 injection	168 days
Fensolvi 6-month	45 mg	1 injection	168 days
Trelstar 1-month	3.75 mg	1 injection	28 days
Trelstar 3-month	11.25 mg	1 injection	84 days
Trelstar 6-month	22.5 mg	1 injection	168 days
Triptodur 6-month	22.5 mg	1 injection	168 days
Supprelin LA	12-month	1 implant	365 days
Synarel	2 mg/mL (200 mcg/spray)	1 bottle (8 mL)	28 days
Zoladex 1-month	3.6 mg	1 implant	28 days
Zoladex 3-month	10.8 mg	1 implant	84 days

CLINICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied

Diagnosis: Gender Identity/Gender Dysphoria				
<u>[niti</u>	al Authorization			
	Select ONE of the following: Member is 18 years of age or older and has a diagnosis of gender dysphoria Member is less than 18 years of age and has a diagnosis of gender dysphoria. Provider please note: parental or legal guardian consent for un-emancipated members is required			
	Provider attests member has the capacity to make informed treatment decisions and has assented to treatment after discussion of the potential benefits and risks			
	 Member has been assessed and diagnosed with gender dysphoria according to DSM-V criteria, by ONE of the following provider types: A licensed mental health provider An endocrinologist A gender dysphoria-informed hormone prescriber, defined as a provider competent in the assessment of gender dysphoria who practices in conjunction with a multidisciplinary gender dysphoria care team 			
	Medication is prescribed by, or in consultation with an endocrinologist or other medical provider experienced in gender dysphoria hormone therapy			
	Provider attests coexisting behavioral health and medical comorbidities or social problems that may interfere with diagnostic procedures or treatment are being appropriately treated and are not causing symptoms of gender dysphoria			
	 Member has experienced puberty development to at least <u>ONE</u> of the following: □ Tanner stage 2 (stage 2 through 4) □ Lab values for Luteinizing Hormone (LH), Follicle Stimulating Hormone (FSH), and the endogenous sex hormones consistent with at least Tanner stage 2 (must submit documentation) If requesting a non-preferred drug, the member has failed <u>ONE</u> of the preferred formulations noted 			
J	above			
Diagnosis: Central Precocious Puberty				
[niti	al Authorization			
	Member is less than 13 years of age			
	Onset of secondary sexual characteristics associated with pubertal pituitary gonadotropin activation, occurring earlier than age 8 for girls and age 9 for boys (submit documentation, progress notes, medical documentation recording physical changes, Tanner staging)			

	Diag	nosis of central precocious puberty is confirmed by <u>ALL</u> the following:		
	□ L	aboratory documentation of pubertal gonadal sex steroid level		
		abertal luteinizing hormone response simulation by native GnRH [Laboratory documentation emonstrating basal LH (>0.3 IU/L), and peak stimulated LH (>4-6 IU/L)]		
	\Box X	-Ray results of the estimated bone age of the non-dominant wrist and hand greater than 2 standard eviations beyond chronological age (submit laboratory and x-ray documentation)		
	tumo	nor has been ruled out by lab tests such as diagnostic imaging of the brain (to rule out intracranial r), pelvic/testicular/adrenal ultrasound (to rule out steroid secreting tumors), and human chorionic lotropin levels (to rule out a chorionic gonadotropin secreting tumor)		
	Medi	cation will NOT be used in combination with growth hormone therapy		
	-	uantity (dose) requested is in accordance with FDA-approved labeling, and if applicable or sary, age and weight conditions are met		
	If req	uesting a non-preferred drug, the member has failed ONE of the preferred formulations noted		
	Diagno	osis: Gynecological Indications		
Init	ial Au	<u>thorization</u>		
	Mem	ber is 18 years of age or older		
	☐ If requesting a non-preferred drug, the member has failed <u>ONE</u> of the preferred formulations noted above			
	Treat	ment is being prescribed by or in consultation with a specialist in obstetrics/gynecology		
Select	ONE	of the following indications for use:		
	symp	UTERINE LEIOMYOMATA, FIBROIDS (requires chart notes documenting tomology/pelvic exam, transvaginal ultrasonography, sono-hysterography): Iember is premenopausal		
		lember has uterine leiomyomas (fibroids)		
		lember is using for the management of heavy menstrual bleeding		
	□ M	lember does <u>NOT</u> have any contraindications to therapy including osteoporosis, history of rombotic or thromboembolic disorders, uncontrolled hypertension, or hepatic impairment/disease		
	□ M	lember has history of inadequate response to the following therapies, tried for at least three (3) onths each (must submit chart note documentation of all therapy failures):		
		tranexamic acid 650 mg		
		<u>OR</u>		
		Member has had surgery for uterine fibroids (i.e., ablation, myomectomy) and has persistent symptoms (must submit documentation of date/type of surgery or procedure)		

		The quantity (dose) and administration frequency requested is in accordance with FDA-approved labeling, and if applicable or necessary, age and weight conditions are met		
	FOR ENDOMETRIOSIS (requires chart notes detailing and recording treatment plan and/or symptomology of chronic pelvic pain (defined as noncyclical pain lasting 6 or more months that localizes to the anatomic pelvis, anterior abdominal wall at or below the umbilicus, the lumbosacral back, or the buttocks, and is of sufficient severity to cause functional disability or lead to medical care amenorrhea, preoperative ablation treatment):			
		Member is premenopausal		
		Member has a diagnosis of moderate to severe pain associated with endometriosis		
		Diagnosis of endometriosis has been confirmed by direct visualization during surgery and/or histology		
		Member does <u>NOT</u> have any contraindications to therapy including osteoporosis, severe hepatic impairment/disease, or concomitant use of hormonal contraceptives		
		Member has history of inadequate response to the following therapies, tried for at least three (3) months each (must submit chart note documentation of all therapy failures):		
		□ NSAIDs (non-steroidal anti-inflammatory drugs		
		☐ Combination (estrogen/progesterone) oral contraceptive		
		□ Progestins		
		<u>OR</u>		
		☐ Member has had surgical ablation to prevent recurrence		
		The quantity (dose) and administration frequency requested is in accordance with FDA-approved labeling, and if applicable or necessary, age and weight conditions are met		
REA	UT	THORIZATION CRITERIA		
	Me	ember continues to meet all initial criteria		
	_	nosis: Central Precocious Puberty (Please submit chart notes and other supporting nents)		
	Me	ember is <u>NOT</u> over the age of 13		
	Member has experienced disease response as indicated by lack of progression or stabilization of secondary sexual characteristics, decrease in growth velocity and bone age advancement, and improvement in final height prediction			
		ember has experienced an absence of unacceptable toxicity from the drug (e.g., convulsions, velopment or worsening of psychiatric symptoms)		

	Diagnosis: Gynecological Indications (Please submit chart notes and other supporting locuments)			
	Total duration of therapy (initial plus re-treatment for symptom recurrence) should not exceed 12 months, AND will be used in combination with add-back therapy, unless clinically contraindicated			
	<u>OR</u>			
	The member's medical history and medical condition's current status requires longer treatment duration than otherwise recommended in published compendia/FDA labeling.			
	Please provide an explanation along with any pertinent progress notes of medical condition including recorded recurrence of symptoms, procedure/exam results:			
	AND			
	Prescriber will order and review a bone density assessment prior to re-treatment			
N #	1' 4' 1 ' ' ' 1 11 C ' 14 DI			
vie	Medication being provided by Specialty Pharmacy – Proprium Rx			

**Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. **

*Previous therapies will be verified through pharmacy paid claims or submitted chart notes. *