	MEDICARE FORM Lupron Depot <sup>®</sup> (leuprolide acetate for depot suspension) Medication Precertification Request Page 1 of 3 (All fields must be completed and legible for precertification review.) reatment: Start date// tion of therapy, Date of last treatment//					For Ohio MMP: FAX: 1-855-734-9389 PHONE: 1-855-364-0974 For other lines of business: Please use other form. Note: Lupron Depot is non- preferred. The preferred product is Eligard. Firmagon is also a preferred product.	
Precertification Requested By	:		Phor	ne:	Fax	:	
A. PATIENT INFORMATION							
First Name:		Last Name:	•		DOB:	I	
Address:			City:		State:	ZIP:	
Home Phone:	Work Phone:		Cell Phone:		Email:		
Patient Current Weight: Ibs	-	ent Height: inche	es or <u> </u>	Allergies:			
<b>B. INSURANCE INFORMATION</b>							
Aetna Member ID #:		Does patient have oth					
Group #: Insured:		If yes, provide ID#: Insured:		Carrier Name:			
		IVIO	edicald: 📋 Yes	☐ No If yes, pro	vide ID #:		
C. PRESCRIBER INFORMATIO First Name:	N	Last Name:		(Check (	)ne)· □ M D	. 🗌 D.O. 🗌 N.P. 🗌 P. <i>I</i>	
Address:		Last Name.	City:	(Onech e	State:	ZIP:	
Phone: Fax	x:	St Lic #:	NPI #:	DEA #:	0.0.101	UPIN:	
Provider Email:		Office Contact Name			Phone:	-	
Agency Name: Administration code(s) (CPT):	Phone:		_	y Pharmacy			
Address:	01-1-	210					
City: Phone:							
TIN:			NPI:				
NPI:			_				
E. PRODUCT INFORMATION							
Request is for: Lupron Depot (I	euprolide acetate fo	r depot suspension) D	)ose:	F	requency:		
F. DIAGNOSIS INFORMATION	Please indicate prim	ary ICD code and spec	ify any other whe	ere applicable.			
Primary ICD Code:		Secondary ICD Cod	e:	Other	ICD Code:		
Yes 🗌 No	ocumentation require r Lupron Depot-PED? upron Depot-PED form sex cord-stromal turn of dose is being require d drug being prescribe Is the patient undergo Will the patient receive unner Stage of puberty umors	ed for all requests): n for this request. nors, prostate cancer, re- ested: 3.75 mg 7 d for pubertal hormonal so bing gender transition? ve the requested drug con- the patient has reached:	ecurrent salivary .5 mg 11.25 suppression in an ncomitantly with g Stage I Sta	gland tumors india mg □ 22.5 mg □ adolescent patient? ender-affirming horn age II □ Stage III [	cations only: ] 30 mg	-	

For Ohio MMP:



## **MEDICARE FORM**

## Lupron Depot<sup>®</sup> (leuprolide acetate for depot suspension) Medication Precertification Request Page 2 of 3

(All fields must be completed and legible for precertification review.)

For Ohio MMP: FAX: 1-855-734-9389 PHONE: 1-855-364-0974

For other lines of business: Please use other form.

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Patient First Name       Patient Last Name       Patient Phone       Patient DOB         G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.       Patient salivary gland tumors							
Recurrent salivary gland tumors							
☐ Yes ☐ No Is the tumor androgen receptor positive?							
For breast cancer, endometriosis, ovarian cancer, preservation of ovarian function, recurrent menstrual related attacks in acute porphyria or uterine							
leiomyomata (fibroids) indication only:							
Please select which Lupron Depot dose is being requested:  3.75 mg 11.25 mg							
Breast cancer							
Please indicate the patient's hormone receptor (HR) status: HR-positive HR-negative Unknown							
Ovarian cancer Please select: Epithelial ovarian cancer Fallopian tube cancer Primary peritoneal cancer Malignant sex cord-stromal tumor							
□ Preservation of ovarian function							
☐ Yes ☐ No Is the patient premenopausal and undergoing chemotherapy?							
Prevention of recurrent menstrual related attacks in acute porphyria							
☐ Yes ☐ No Is the requested drug being requested to prevent recurrent menstrual related attacks in acute porphyria?							
Yes No Is the requested drug being prescribed by, or in consultation with, a physician experienced in the management of porphyrias?							
Uterine leiomyomata (fibroids)							
Yes No Does the patient have a diagnosis of anemia (for example, Hct less than or equal to 30% and/or Hgb less than or equal to 10 g/dL)?							
$\square$ Yes $\square$ No Will the requested drug be used prior to surgery for uterine fibroids?							
For Continuation Requests (clinical documentation required for all requests):							
For gender dysphoria, malignant sex cord-stromal tumors, prostate cancer, recurrent salivary gland tumors continuation requests only:							
Please select which Lupron Depot dose is being requested: 🗌 3.75 mg 📋 7.5 mg 📋 11.25 mg 📋 22.5 mg 🔲 30 mg 🔲 45 mg							
Gender dysphoria							
Yes No Is the requested drug being prescribed for pubertal hormonal suppression in an adolescent patient?							
$\longrightarrow$ Yes $\square$ No Is the patient undergoing gender transition?							
Yes No Will the patient receive the requested drug concomitantly with gender-affirming hormones?							
└────────────────────────────────────	n						
☐ Malignant sex cord-stromal tumors							
Yes No Has the patient experienced an unacceptable toxicity or disease progression while receiving the requested drug?							
□ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □							
☐ Yes ☐ No Has the patient had prior therapy with Lupron Depot within the last 365 days?							
🗌 Yes 🔲 No Has the patient experienced clinical benefit while receiving the requested drug (e.g., serum testosterone less than 50ng/dl)?							
Yes No Has the patient experienced an unacceptable toxicity while receiving the requested drug?							
Recurrent salivary gland tumors							
Yes No Has the patient experienced an unacceptable toxicity or disease progression while receiving the requested drug?							

Continued on next page



## **MEDICARE FORM**

## Lupron Depot<sup>®</sup> (leuprolide acetate for depot suspension) Medication Precertification Request Page 3 of 3

(All fields must be completed and legible for precertification review.)

For Ohio MMP: FAX: 1-855-734-9389 PHONE: 1-855-364-0974

For other lines of business: Please use other form.

Note: Lupron Depot is nonpreferred. The preferred product is Eligard. Firmagon is also a preferred product.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.								
For breast cancer, endometriosis, ovarian cancer, preservation of ovarian function, recurrent menstrual related attacks in acute porphyria or uterine								
fibroids continuation requests only:								
Please select Lupron Depot dose for the following indications: 🗌 3.75 mg 🔲 11.25 mg								
Breast cancer								
Please indicate the patient's hormone receptor (HR) status: 🗌 HR-positive 🔲 HR-negative 🔲 Unknown								
Yes No Has the patient experienced clinical benefit while receiving the requested drug?								
☐ Yes ☐ No Has the patient experienced an unacceptable toxicity while receiving the requested drug?								
Endometriosis								
Yes No Has the patient received previous therapy with the requested medication or Lupaneta Pack?								
└────────────────────────────────────								
Yes No Is the patient's bone mineral density within normal limits?								
How long has the patient received previous therapy with the requested drug and Lupaneta Pack? months								
Ovarian cancer								
Please select: 🗌 Epithelial ovarian cancer 🔲 Fallopian tube cancer 🔲 Primary peritoneal cancer 🔲 Malignant sex cord-stromal tumor								
Yes INo Has the patient experienced clinical benefit while receiving the requested drug?								
Yes No Has the patient experienced an unacceptable toxicity while receiving the requested drug?								
Preservation of ovarian function								
_ Yes □ No Is the patient premenopaus								
Prevention of recurrent menstrual related								
Yes No Is the requested medication being requested to prevent recurrent menstrual related attacks in acute porphyria?								
🗌 Yes 🔲 No Is the requested medication being prescribed by, or in consultation with, a physician experienced in the management of porphyrias?								
Uterine leiomyomata (fibroids)								
☐ Yes ☐ No Has the patient received previous therapy with the requested drug or Lupaneta Pack?								
→ Yes No Does the patient have a diagnosis of anemia (for example, Hct less than or equal to 30% and/or Hgb less than or equal to 10g/dL)?								
How long has the patient received previous therapy with the requested drug and Lupaneta Pack? months								
└────────────────────────────────────	patient have a diagnosis of anemia (for ex 10g/dL)?	ample, Hct less than or equal to	30% and/or Hgb less than or					
└───> □ Yes [	No Will the requested drug be used pr	ior to surgery for uterine fibroids?						
H. ACKNOWLEDGEMENT								
Request Completed By (Signature Require	ed):		Date: / /					
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive								
any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent								

insurance act, which is a crime and subjects such person to criminal and civil penalties.

The plan may request additional information or clarification, if needed, to evaluate requests.