

## **MEDICARE FORM**

## Zoladex® (goserelin acetate) Medication Precertification Request

Page 1 of 2

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

For Michigan MMP: FAX: 1-844-241-2495 PHONE: 1-855-676-5772 For other lines of business:

Please use other form.

Note: Zoladex is non-preferred. The preferred product is Eligard. Firmagon is also a preferred product.

	rt of treatment: Start date _				•		
	ntinuation of therapy, Date o	of last treatment			<b></b>		
Precertification Request			Phone	::	Fax:		
A. PATIENT INFORMATION	ON						
First Name:		Last Name:	Т		DOB:	1	
Address:	<del></del>		City:		State:	ZIP:	
Home Phone:	Work Phone:		Cell Phone:		Email:		
Patient Current Weight:	lbs orkgs Patie	nt Height: inches	orcms	Allergies:			
B. INSURANCE INFORM	ATION						
					☐ Yes ☐ No		
	Group #:		If yes, provide ID#: Carrier		r Name:		
Insured:		Insured:					
Medicare: Yes No		Me	edicaid: ∐ Yes	☐ No If yes, pro	ovide ID #:		
C. PRESCRIBER INFORM	MATION	Last Name:		(Chook C			
First Name:		Last Name:	0:4	(Crieck C		☐ D.O. ☐ N.P. ☐ P.A.	
Address:	<u></u>	T "	City:		State:	ZIP:	
Phone:	Fax:	St Lic #:	NPI #:	DEA #:	T	UPIN:	
Provider Email:		Office Contact Name:	:		Phone:		
Specialty (Check one):	Oncologist	logist 🗌 Other:					
Address:	Phone:  CPT): State: Z  Fax: PIN:	ZIP:	Specialty Name: Address: City: Phone: TIN:		Other: State: Fax: PIN:	ZIP:	
	(goserelin acetate) Dose:		Frequenc	:v:			
_ =	TION - Please indicate prima						
Primary ICD Code:		Secondary ICD Code			r ICD Code:		
	ION - Required clinical inform	<del>-</del>	<u> </u>	in its entirety for all precertification requests.			
For Initiation Requests (clinical documentation required for all requests):  For Zoladex 3.6 mg requests only:  Breast cancer  Please indicate the patient's hormone receptor (HR) status: HR-positive HR-negative Unknown  Chronic anovulatory uterine bleeding  Yes No Will the requested medication be used as an endometrial thinning agent prior to endometrial ablation for dysfunctional uterine bleeding?  We will the requested medication be used for treatment of chronic anovulatory uterine bleeding in a patient with severe anemia?  Dysfunctional uterine bleeding  Yes No Will the requested medication be used as an endometrial thinning agent prior to endometrial ablation for dysfunctional uterine bleeding?  Yes No Will the requested medication be used for treatment of chronic anovulatory uterine bleeding in a patient with severe anemia?							
Please indicate how many months has the patient already received the requested medication for this indication: 6 months or greater Less than 6 months							



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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.								
☐ Gender dysphoria								
Yes No Is the requested medication being prescribed for pubertal suppression in an adolescent patient?								
<ul> <li>Yes ☐ No Is the patient undergoing gender transition?</li> <li>Yes ☐ No Will the patient receive the requested medication concomitantly with gender affirming hormones?</li> </ul>								
Please indicate the Tanner Stage of puberty the patient has reached: Stage I Stage II Stage II Stage IV Stage IV Unknown								
☐ Preservation of ovarian function								
Yes No Is the patient premenopausal and undergoing chemotherapy?								
<ul> <li>☐ Prevention of recurrent menstrual related attacks in acute porphyria</li> <li>☐ Yes ☐ No Is the requested medication being requested to prevent recurrent menstrual related attacks in acute porphyria?</li> </ul>								
Yes No Is the requested medication being requested to prevent recurrent mensitual 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?								
Prostate cancer								
Note: Zoladex is non-preferred. The preferred product is Eligard. Firmagon is also a preferred product.								
☐ Yes ☐ No Has the patient had a trial and failure, intolerance, or contraindication to Eligard?								
Please explain if there are any other medical reason(s) that the patient cannot use Eligard when indicated for the patient's diagnosis?								
Uterine leiomyomata (fibroids)	modication has given prior to aurgeny?							
☐ Yes ☐ No Will the requested medication be given prior to surgery?  For Zoladex 10.8 mg requests only:								
☐ Breast cancer								
<del>-</del>	recentor (HR) status: \( \Price \text{HR-positive} \)	HR-negative D Unknown						
Please indicate the patient's hormone receptor (HR) status: HR-positive HR-negative Unknown								
☐ Gender dysphoria ☐ Yes ☐ No Is the requested medication being prescribed for pubertal suppression in an adolescent patient?								
	s the patient undergoing gender transitio		•					
	Will the patient receive the requested me							
Please indicate the Tanner Stage of puberty the patient has reached: Stage I Stage II Stage III Stage IV Stage V Unknown								
☐ Prostate cancer         ☐ Yes       ☐ No       Has the patient had an ineffective response, contraindication, or intolerance to Eligard?								
Yes No Has the patient had an ineffective response, contraindication, or intolerance to Firmagon?								
For Continuation Requests (clinical of	locumentation required for all request	s):						
☐ Breast cancer		<u> </u>						
☐ 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?								
☐ Gender dysphoria ☐ Yes ☐ No Is the requested medication being prescribed for pubertal suppression in an adolescent patient?								
Yes No is the requested medication being prescribed for pubertal suppression in an adolescent patient?  Yes No is the patient undergoing gender transition?								
Yes No Will the patient receive the requested medication concomitantly with gender affirming hormones?								
Please indicate the Tanner Stage of puberty the patient has reached:  Stage I  Stage II  Stage III  Stage IV  Stage V  Unknown								
Preservation of ovarian function								
☐ Yes ☐ No Is the patient premenopausal and still undergoing chemotherapy? ☐ Prevention of recurrent menstrual related attacks in acute porphyria								
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?								
☐ Prostate cancer								
•	d prior therapy with Zoladex within the la	-	/					
☐ Yes ☐ No Has the patient experienced clinical benefit to therapy while receiving the requested drug (e.g., serum testosterone less than 50 ng/dl)?								
☐ Yes ☐ No Has the patient experienced an unacceptable toxicity while receiving the requested drug?								
H. ACKNOWLEDGEMENT								
	Paguirod):		Date: / /					
Request Completed By (Signature								
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.