

1230 US Highway 11

Gouverneur, NY 13642

Phone: 1-877-635-9545

Prior Authorization Fax: 1-844-712-8129

Testosterone Enanthate Prior Authorization Request Form (Page 1 of 3)

Member Information (required)			Provider Information (required)			
Member Name:			Provider Name:			
Insurance ID#:			NPI#: Specialty:			
Date of Birth:			Office Phone:			
Street Address:			Office Fax:			
City:	State:	Zip:	Office Street Address:			
Phone:			City: State: Zip:		Zip:	
		Medication Info	ormation (required)			
Medication Name:			Strength: Dosage Form:		orm:	
☐ Check if requesting brand			Directions for Use:			
☐ Check if request is t	for continuation of the	rapy				
		Clinical Inforr	nation (required)			
Select the diagnosis below: Delayed puberty Female-to-male transsexual -gender identity disorder Hypogonadotropic hypogonadism (congenital or acquired) Inoperable breast cancer in women Primary hypogonadism (congenital or acquired) Other diagnosis: ICD-10 Code(s):						
For delayed puberty, answer the following: Was the patient a male at birth? Yes No						
For female-to-male transsexual - gender identity disorder, answer the following: Is the patient using hormones to change physical characteristics?						
For inoperable breast cancer in women, answer the following: Is the medication being used for palliative treatment of inoperable breast cancer? ☐ Yes ☐ No Was the patient a female at birth? ☐ Yes ☐ No						



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For male hypoge	onadism, answer the followi	ng:					
Was the patient a	a male at birth? Yes No						
Does the patient range for the lab?		total testosterone levels less than 2	280 ng/dL (<9.7 nmol/L) or less than the reference				
		I testosterone levels and the referen					
Total testostero	one level:	Reference range:	Date taken:				
Total testostero	one level:	_Reference range:	Date taken:				
Does the patient obesity)? ☐ Yes		se altered sex-hormone binding glob	bulin (SHBG) (e.g., thyroid disorder, diabetes,				
	Does the patient have one pre-treatment calculated free or bioavailable testosterone level less than 5 ng/dL (< 0.17 nmol/L) or less than the reference range for the lab? Yes No						
Please document	the pre-treatment calculated	free or bioavailable testosterone leve	el and the reference range for the lab:				
Free testostero	ne level:	_Reference range:	Date taken:				
□ Bilateral orcl□ Panhypopitu	itarism	nadism (e.g., congenital anorchia, k	Klinefelter's syndrome)				
Reauthorization	:						
If this is a reauth	norization request for gende	r identity disorder or male hypogo	onadism, answer the following questions:				
	apy, or 12 months for patients		the past 6 months for patients new to t is within or below the normal limits of the reporting				
Please document	the serum total testosterone	evel and the reference ranges for th	ne lab:				
Current total te	stosterone level:	Reference range:	Date taken:				
testosterone there		continuing testosterone therapy that	the past 6 months for patients new to t is outside of upper limits of normal for the reporting				
Does the patient obesity)? ☐ Yes		se altered sex-hormone binding glob	bulin (SHBG) (e.g., thyroid disorder, diabetes,				
	erapy, or 12 months for patier		drawn from within the past 6 months for patients new hat is within or below the normal limits of the				
		rone level and the reference ranges					
Current free/bid	pavailable testosterone level:_	Reference range:	Date taken:				
to testosterone th		nts continuing testosterone therapy t	drawn from within the past 6 months for patients new hat is outside of upper limits of normal for the				
Are there any other this review?	er comments, diagnoses, sympt	oms, medications tried or failed, and/	or any other information the physician feels is important to				
			_				
Please note:	•	ss all required information is received.					
	Please fax this form to 1-844-7	12-8129 to initiate a prior authorizatio	n review for the member and medication above.				
	Please note: plan benefits may	/ limit or exclude coverage of specific	medications including those requested on this form.				



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Provider/Representative (and Title): Date:							
PROACT INTERNAL USE ONLY:							
Clinical Review Decision							
	Approved, through						
	Denied (documentation attached, if necessary)						
Tracking:							
1st Attemp	ot		2 nd Attempt		Letter Mailed:		

I certify, to the best of my knowledge, the statements and information provided on this form are factual and correct.