## **TESTOSTERONES PRIOR AUTHORIZATION REQUEST FORM**



OptumRx
P.O. Box 25184
Santa Ana, CA, 92799
Phone: (800) 310-6826 Fax: (866) 940-7328



Note: This form must be complete **All sections		•	will be returned**
Patient's Medicaid #		Date of Birth	//
Patient's Name		Prescriber's Nan	ne
Prescriber's IN License #		Specialty	
Prescriber's NPI#		Prescriber's Sigr	nature
Return Fax #		Return Phone #	
Check box if requesting retro-active PA	A	Date(s) of service retro-active eligit	e requested for oility (if applicable):
lote: Submit PA requests for retroactive ligibility timelines) with dates of service pf service 30 calendar days or less and g	orior to 30 calendar d		determination, but within established eparately from current PA requests (dates
Requested Medication	Strength	Quantity	Dosage Regimen
DEPO-TESTOSTERONE, TESTOS Initial Authorization:	TERONE CYPION	ATE	
Initial Authorization:  1. Please select one of the following  Member has a diagnosis of o	: delayed puberty one level ≤ 350 ng, s none of the follow ber assigned male	/dL within the past ving contraindication at birth	

If <b>no</b> , please specify contraindication and medical rationale for use:
in no, please specify contraindication and medical rationale for use.
TESTOSTERONE ENANTHATE
Initial Authorization:
1. Please select one of the following:
<ul> <li>Member has a diagnosis of delayed puberty</li> <li>Has the member had a previous trial and failure of ALL preferred injectable testosterone agents, as</li> </ul>
confirmed by claims history, chart documentation, or provider attestation including dates of trial
(reference PA criteria)? ☐ Yes ☐ No
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:
<ul> <li>Member has a total testosterone level ≤ 350 ng/dL within the past 3 months (Documentation is required)</li> <li>Has the member had a previous trial and failure of ALL preferred injectable testosterone agents (reference PA criteria)? ☐ Yes ☐ No</li> </ul>
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:
<ul> <li>☐ Member needs medication for palliative treatment of metastatic breast cancer</li> <li>2. For ALL indications: <ul> <li>Provider attests that member has none of the following contraindications to therapy: ☐ Yes ☐ No</li> <li>Breast cancer in a member assigned male at birth</li> <li>Pregnancy</li> <li>Prostate cancer</li> </ul> </li> </ul>
If <b>no</b> , please specify contraindication and medical rationale for use:
Note: If member has had history with injectable/topical product within the past 120 days, confirmed by claims history, and are switching formulations to nonpreferred injectable formulation, reauthorization criteria will apply.
Reauthorization:
1. Total testosterone level is $\leq$ 1000 ng/dL within the past 6 months (Documentation is required) $\square$ Yes $\square$ No
2. Has the member had a previous trial and failure of at least ONE preferred injectable testosterone agent, as confirmed by claims history, chart documentation, or provider attestation including dates of trial (not required for palliative treatment of breast cancer) [reference PA criteria]?   Yes  No
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:
3. Provider attests that member remains a candidate for treatment, indicating that they have not developed any of the contraindication(s) listed under initial authorization above $\square$ Yes $\square$ No

If <b>no</b> , please specify contraindication and medical rationale for use:
——————————————————————————————————————
AVEED TEOTODEL DELLET VVOOTED
AVEED, TESTOPEL PELLET, XYSOTED Initial Authorization:
Please select one of the following:
$\square$ Member has a diagnosis of delayed puberty
<ul> <li>Has the member had a previous trial and failure of ALL preferred injectable testosterone agents, as confirmed by claims history, chart documentation, or provider attestation including dates of trial (reference PA criteria)?   Yes  No</li> </ul>
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:
<ul> <li>Member has a total testosterone level ≤ 350 ng/dL within the past 3 months (Documentation is required)</li> <li>Has the member had a previous trial and failure of ALL preferred injectable testosterone agents (reference PA criteria)? ☐ Yes ☐ No</li> </ul>
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:
<ul> <li>2. For ALL indications: Provider attests that member has none of the following contraindications to therapy: □ Yes □ No <ul> <li>Breast cancer in a member assigned male at birth</li> <li>Hypogonadal conditions not associated with structural or genetic etiologies (Xyosted ONLY)</li> <li>Pregnancy</li> <li>Prostate cancer</li> </ul> </li> </ul>
If <b>no</b> , please specify contraindication and medical rationale for use:
Note: If member has had history with injectable/topical product within the past 120 days, confirmed by claims history, and are switching formulations to nonpreferred injectable formulation, reauthorization criteria will apply
Reauthorization:  1. Total testosterone level is ≤1000 ng/dL within the past 6 months (Documentation is required) ☐ Yes ☐ No
2. Has the member had a previous trial and failure of at least ONE preferred injectable testosterone agent, as confirmed by claims history, chart documentation, or provider attestation including dates of trial (reference PA criteria)?   Yes  No
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:
3. Provider attests that member remains a candidate for treatment, indicating that they have not developed any of the contraindication(s) listed under initial authorization above ☐ Yes ☐ No

If <b>no</b> , please specify contraindication and medical rationale for use:
ANDRODERM, TESTOSTERONE 1% (25 MG)/ 2.5 GM GEL PACKETS, TESTOSTERONE 1% (12.5 MG)/ACT GEL PUMP, TESTOSTERONE 1.62% (20.25 MG)/ACT METERED PUMP GEL, TESTIM 1% (50 MG)/5 GM GEL TUBES
Initial Authorization:  1. Please select one of the following:
<ul> <li>Member is 16 years of age or older, has a total testosterone level ≤ 350 ng/dL within the past 3 months (Documentation is required), and is requesting to use topical testosterone within the established quantity limits □ Yes □ No</li> </ul>
IIIIIIts   Yes   No
Requested dose:
<ul> <li>Member is 16 years of age or older, has a total testosterone level ≤ 400 ng/dL while on topical testosterone therapy (Documentation is required) and is requesting to exceed established quantity limits</li> <li>Yes □ No</li> </ul>
Requested dose:
Member has utilized ≥ 14 days of topical testosterone therapy: ☐ Yes ☐ No
Name of medication:
Dose:
Start and End date:
If <b>no</b> , please provide medical justification as to why member is requesting a dose beyond established quantity limits:
2. For <b>ALL</b> indications:  Provider attests that member has none of the following contraindications to therapy: □ Yes □ No  • Breast cancer in a member assigned male at birth  • Pregnancy  • Prostate cancer
If <b>no</b> , please specify contraindication and medical rationale for use:
Note: If member has had history with injectable/topical product within the past 120 days, confirmed by claims history, and are switching formulations to nonpreferred injectable formulation, reauthorization criteria will apply
Reauthorization:  1. Total testosterone level is ≤ 1000 ng/dL within the past 6 months (Documentation is required) ☐ Yes ☐ No
2. Provider attests that member remains a candidate for treatment, indicating that they have not developed any of the contraindication(s) listed under initial authorization above $\square$ Yes $\square$ No

Note: dos	se requested for reauthorization should not exceed established quantity limits unless member
	ly has been approved to exceed the established quantity limits
Reque	sted dose:
(40.5 MG TESTOS	O, TESTOSTERONE 1% (50 MG)/5 GM GEL PACKETS/TUBES, TESTOSTERONE 1.62% b)/2.5 GM GEL PACKETS, TESTOSTERONE 1.62% (20.25 MG)/1.25 GM GEL PACKETS, TERONE 2% (10 MG)/ACT METERED PUMP, TESTOSTERONE 30 MG/ACT SOLUTION, (O 1% (50 MG)/5 GM GEL PACKETS, VOGELXO 1% (12.5 MG)/ACT GEL PUMP
	thorization:
	select one of the following: ember is 16 years of age or older, has a total testosterone level ≤ 350 ng/dL within the past 3 months
(Do	cumentation is required), and is requesting to use topical testosterone within the established quantity lits $\square$ Yes $\square$ No
	Requested dose:
tes	ember is 16 years of age or older, has a total testosterone level ≤ 400 ng/dL while on topical stosterone therapy (Documentation is required) and is requesting to exceed established quantity limits  Yes □ No
	Requested dose:
	Member has utilized ≥ 14 days of topical testosterone therapy: ☐ Yes ☐ No
	Name of medication:
	Dose:
	Start and End date:
	If <b>no</b> , please provide medical justification as to why member is requesting a dose beyond established quantity limits:
document If r	is trial and failure of ALL preferred topical testosterone agents, as confirmed by claims history, chart ation, or provider attestation including dates of trial (reference PA criteria)   Yes  No  No, please provide medical justification for use of requested agent over ALL preferred topical tosterone agents:
	L indications: ler attests that member has none of the following contraindications to therapy: ☐ Yes ☐ No Breast cancer in a member assigned male at birth Pregnancy Prostate cancer

Reauthorization:  1. Total testosterone level is ≤1000 ng/dL within the past 6 months (Documentation is required) ☐ Yes ☐ No  2. Previous trial and failure of at least ONE preferred topical testosterone agent, as confirmed by claims histochart documentation, or provider attestation including dates of trial (reference PA criteria) ☐ Yes ☐ No  If no, please provide medical justification for use of requested agent over ALL preferred topical testosterone agents:
Previous trial and failure of at least ONE preferred topical testosterone agent, as confirmed by claims historian documentation, or provider attestation including dates of trial (reference PA criteria) $\square$ Yes $\square$ No If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred topical
hart documentation, or provider attestation including dates of trial (reference PA criteria)
. Provider attests that member remains a candidate for treatment, indicating that they have not developed a the contraindication(s) listed under initial authorization above $\square$ Yes $\square$ No
If <b>no</b> , please specify contraindication and medical rationale for use:
ANAZOL:
nitial Authorization (approval up to 6 months):
. Member diagnosis(es):
ote: Approvable diagnoses include angioedema prophylaxis for heredity angioedema, autoimmune hemolytic anemia, disco upus erythematosus, endometriosis, fibrocystic breast disease, myelosclerosis with myeloid metaplasia
. For <b>ALL</b> indications:
Provider attests that member has none of the following contraindications to therapy: $\Box$ Yes $\Box$ No
Active or history of thrombosis or thromboembolic disease
Androgen-dependent tumor
<ul><li>Androgen-dependent tumor</li><li>Cardiac disease</li></ul>
<ul> <li>Androgen-dependent tumor</li> <li>Cardiac disease</li> </ul>
<ul> <li>Androgen-dependent tumor</li> <li>Cardiac disease</li> <li>Porphyria</li> <li>Pregnancy or breast-feeding</li> <li>Severe hepatic disease</li> </ul>
<ul> <li>Androgen-dependent tumor</li> <li>Cardiac disease</li> <li>Porphyria</li> <li>Pregnancy or breast-feeding</li> <li>Severe hepatic disease</li> <li>Severe renal disease</li> </ul>
<ul> <li>Androgen-dependent tumor</li> <li>Cardiac disease</li> <li>Porphyria</li> <li>Pregnancy or breast-feeding</li> <li>Severe hepatic disease</li> </ul>
<ul> <li>Androgen-dependent tumor</li> <li>Cardiac disease</li> <li>Porphyria</li> <li>Pregnancy or breast-feeding</li> <li>Severe hepatic disease</li> <li>Severe renal disease</li> </ul>

adverse events ☐ Yes ☐ No
2. Provider attests that member remains a candidate for treatment, indicating that they have not developed any of the contraindication(s) listed under initial authorization above $\square$ Yes $\square$ No
If <b>no</b> , please specify contraindication and medical rationale for use:
LATENTO (TEGTOSTEDONE UNDEGANIOATE)
JATENZO (TESTOSTERONE UNDECANOATE): Initial Authorization:
<ol> <li>Member is 18 years of age or older and is requesting to use oral testosterone within the established quantity limits ☐ Yes ☐ No</li> </ol>
Requested dose:
2. Member has a diagnosis of hypogonadism with a total testosterone level ≤ 350 ng/dL within the past 3 months (Documentation is required) ☐ Yes ☐ No
3. Previous trial and failure of at least ONE preferred injectable testosterone agent, as confirmed by claims history, chart documentation, or provider attestation including dates of trial (reference PA criteria) $\square$ Yes $\square$ No
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:
<ul> <li>4. For ALL indications: Provider attests that member has none of the following contraindications to therapy:  <ul> <li>Yes</li> <li>No</li> <li>Breast cancer in a member assigned male at birth</li> <li>Hypogonadal conditions not associated with structural or genetic etiologies</li> <li>Pregnancy</li> <li>Prostate cancer</li> </ul> </li> </ul>
If <b>no</b> , please specify contraindication and medical rationale for use:
Reauthorization:
1. Total testosterone level is ≤ 1000 ng/dL within the past 6 months (Documentation is required) ☐ Yes ☐ No
2. Provider attests that member remains a candidate for treatment, indicating that they have not developed any of the contraindication(s) listed under initial authorization above $\square$ Yes $\square$ No
If <b>no</b> , please specify contraindication and medical rationale for use:
3. Previous trial and failure of at least ONE preferred injectable testosterone agent , as confirmed by claims history, chart documentation, or provider attestation including dates of trial (reference PA criteria)  □ Yes □ No
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:

Note: dose requested for reauthorization should not exceed established quantity limits
Requested dose:
METHITEST (METHYLTESTOSTERONE)
Initial Authorization (approval up to 6 months):
<ul> <li>1. Please select one of the following:</li> <li> Member has a diagnosis of cryptorchidism Member has a diagnosis of delayed puberty Member has a diagnosis of hypogonadism (primary or hypogonadotropic) with a total testosterone ≤ 350 ng/dL within the past 3 months (Documentation is required) Member needs medication for palliative treatment of metastatic breast cancer </li> </ul>
·
2. Previous trial and failure of at least ONE preferred injectable testosterone agent, as confirmed by claims history, chart documentation, or provider attestation including dates of trial (reference PA criteria) $\square$ Yes $\square$ No
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:
<del></del>
3. For <b>ALL</b> indications:
<ul> <li>Provider attests that member has none of the following contraindications to therapy: ☐ Yes ☐ No</li> <li>Breast cancer in a member assigned male at birth</li> <li>Pregnancy</li> <li>Prostate cancer</li> </ul>
If <b>no</b> , please specify contraindication and medical rationale for use:
4. Dose requested of methyltestosterone is within the established quantity limits
Requested dose:
Reauthorization (approval up to 6 months):
Please select one of the following:
Member has a diagnosis of hypogonadism and a total testosterone level ≤ 1000 ng/dL within the past 6 months (Documentation is required)
Member has a diagnosis of delayed puberty, palliative treatment of metastatic breast cancer, or cryptorchidism AND prescriber has submitted documentation indicating continued benefit from the medication without significant adverse events:
2. For <b>ALL</b> indications: Provider attests that member remains a candidate for treatment, indicating that they have not developed any of the contraindication(s) listed under initial authorization above ☐ Yes ☐ No
If <b>no</b> , please specify contraindication and medical rationale for use:

3. Previous trial and failure of at least ONE preferred injectable testosterone agent, as confirmed by claims history, chart documentation, or provider attestation including dates of trial (reference PA criteria) $\square$ Yes	
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:	
	_
Note: dose requested for reauthorization should not exceed established quantity limits	
Requested dose:	
TLANDO (TESTOSTERONE UNDECANOATE)	
<ul> <li>Initial Authorization:</li> <li>1. Member is 18 years of age or older and is requesting to use oral testosterone within the established qualimits ☐ Yes ☐ No</li> </ul>	uantity
Requested dose:	
2. Member has a diagnosis of hypogonadism and a total testosterone level ≤ 350 ng/dL within the past 3 m (Documentation is required) ☐ Yes ☐ No	onths
3. Previous trial and failure of at least ONE preferred injectable testosterone agent , as confirmed by claims history, chart documentation, or provider attestation including dates of trial (reference PA criteria) $\square$ Yes	
If <b>no</b> , please provide medical justification for use of requested agent over ALL preferred injectable testosterone agents:	_
4. For <b>ALL</b> indications:	_
Provider attests that member has none of the following contraindications to therapy: ☐ Yes ☐ No  • Breast cancer	
<ul> <li>Hypogonadal conditions not associated with structural or genetic etiologies</li> <li>Pregnancy</li> <li>Prostate cancer</li> </ul>	
If <b>no</b> , please specify contraindication and medical rationale for use:	
	_
Reauthorization:  1. Total testosterone level is ≤ 1000 ng/dL within the past 6 months (Documentation is required) □ Yes □ No	
2. Prescriber attests that member remains a candidate for treatment, indicating that they have not developed of the contraindication(s) listed under initial authorization above $\square$ Yes $\square$ No	∍d any
If <b>no</b> , please specify contraindication and medical rationale for use:	
	_
3. Previous trial and failure of at least ONE preferred injectable testosterone agent , as confirmed by claims	- S
history, chart documentation, or provider attestation including dates of trial (reference PA criteria) $\square$ Yes $\square$	

Note:	dose requested for reauthorization should not exceed established quantity limits
Red	uested dose:
UNDE	CATREX (TESTOSTERONE UNDECANOATE):
1. Mer	Authorization: nber is 18 years of age or older and is requesting to use oral testosterone within the established quantity its $\Box$ Yes $\Box$ No
Red	uested dose:
	nber has a diagnosis of hypogonadism with a total testosterone level $\leq$ 350 ng/dL within the past 3 months sumentation is required) $\square$ Yes $\square$ No
	vious trial and failure of BOTH Jatenzo AND Tlando, as confirmed by claims history, chart documentation, vider attestation including dates of trial (reference PA criteria) $\square$ Yes $\square$ No
	If <b>no</b> , please provide medical justification for use of requested agent over Jatenzo AND Tlando:
	ALL indications:
	ovider attests that member has none of the following contraindications to therapy: $\Box$ Yes $\Box$ No
	<ul> <li>vider attests that member has none of the following contraindications to therapy:  <ul> <li>Yes</li> <li>No</li> </ul> </li> <li>Breast cancer in a member assigned male at birth</li> <li>Hypogonadal conditions not associated with structural or genetic etiologies</li> <li>Pregnancy</li> </ul>
	<ul> <li>ovider attests that member has none of the following contraindications to therapy:  <ul> <li>Yes</li> <li>No</li> </ul> </li> <li>Breast cancer in a member assigned male at birth</li> <li>Hypogonadal conditions not associated with structural or genetic etiologies</li> </ul>
	<ul> <li>vider attests that member has none of the following contraindications to therapy:  <ul> <li>Yes</li> <li>No</li> </ul> </li> <li>Breast cancer in a member assigned male at birth</li> <li>Hypogonadal conditions not associated with structural or genetic etiologies</li> <li>Pregnancy</li> </ul>
Pr	byvider attests that member has none of the following contraindications to therapy:
Pro	<ul> <li>vider attests that member has none of the following contraindications to therapy:  <ul> <li>Yes</li> <li>No</li> </ul> </li> <li>Breast cancer in a member assigned male at birth</li> <li>Hypogonadal conditions not associated with structural or genetic etiologies</li> <li>Pregnancy</li> <li>Prostate cancer</li> </ul>
Reaut 1. Tota 2. Pro	byider attests that member has none of the following contraindications to therapy:
Reaut 1. Tota 2. Pro	• Breast cancer in a member assigned male at birth • Hypogonadal conditions not associated with structural or genetic etiologies • Pregnancy • Prostate cancer  If no, please specify contraindication and medical rationale for use:    horization:   all testosterone level is ≤ 1000 ng/dL within the past 6 months (Documentation is required)
Reaut 1. Tota 2. Pro the	ovider attests that member has none of the following contraindications to therapy: ☐ Yes ☐ No  ■ Breast cancer in a member assigned male at birth  ■ Hypogonadal conditions not associated with structural or genetic etiologies  ■ Pregnancy  ■ Prostate cancer  If no, please specify contraindication and medical rationale for use:  ———————————————————————————————————
Reaut 1. Tota 2. Prothe 3. Pretor prov	ovider attests that member has none of the following contraindications to therapy: ☐ Yes ☐ No  ■ Breast cancer in a member assigned male at birth  ■ Hypogonadal conditions not associated with structural or genetic etiologies  ■ Pregnancy  ■ Prostate cancer  If no, please specify contraindication and medical rationale for use:    If no   Prostate cancer   Prostate cance

Note: dose requested for reauthorization should not exceed established quantity limits	
Requested dose:	

## CONFIDENTIAL INFORMATION

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