

## BlueShield. TESTOSTERONE (BUCCAL/NASAL/ORAL) Federal Employee Program. PRIOR APPROVAL REQUEST

Additional information is required to process your claim for prescription drugs. Please complete the patient portion, and have the prescribing physician complete the physician portion and submit this completed form.

Send completed form to: Service Benefit Plan Prior Approval P.O. Box 52080 MC 139 Phoenix, AZ 85072-2080 Attn. Clinical Services Fax: 1-877-378-4727

Patient Information (required)			Provider Information (required)					
Date:			Provider Name:					
Patient Name:			Specialty:		NPI:			
Date of Birth:	Sex: □Male	□Female	Office Phone:		Office Fax:			
Street Address:			Office Street Address:					
City:	State:	Zip:	City:	Sta	ite:	Zip:		
Patient ID: R	1 1		Physician Signature:					
PHYSICIAN COMPLETES								
Testosterone (Buccal/Nasal/Oral)								
<b>NOTE</b> : Form must be completed in its <b>entirety</b> for processing								
Please select the product being requested:								
□ Jatenzo 158mg capsule □ Jatenzo 198mg capsule □ Jatenzo 237mg capsule □ Jatenzo 237mg capsule □ Mothyltestastaror			□Natesto nasal gel □Striant buccal system capsule □Tlando					
□Jatenzo 237mg capsule		thyltestostero		Tlando				
*Check www.fepblue.org/formulary to o ***Non-covered branded medication				eption pr	ocess			
		_	,, ,, ,	F				
s this request for brand or generic?  Generic  Generic								
. How many (bottles/capsules/tab	lets) will the patie	ent need for a 9	90 day supply?	_ every 9	0 days			
<ol> <li>Will this medication be used in a *If YES, please specify the m</li> </ol>		any other form	n of testosterone? □Yes*	□No				
3. Is the patient being treated for ger <b>QYES</b> : Is the patient undergoin		-		sformatio	n, or sex chang	ge? Answer below:		
<b>□NO</b> : Please answer the follow	ing questions:							
a. Is the patient assigned	female or male at	birth? □Fem	ale <u>OR</u>					
b. What is the patient's d	•							
Delay in sexual deve		•	41	1 1. · ·		dia amandai		
1. Will the patient evidence? □Ye		nand and wris	t be assessed every 6 mont	ns as dete	ermined by ra	aiograpnic		
ii. Will the patient	's liver functions	tests be monito	ored every 6 months? $\Box$ Y	es 🗆 N	О			
iii. Will the patient's hematocrit levels be monitored every 6 months? □Yes □No								
	been on testoster excluding sample		n any dosage form (injection No	on, topica	l, oral, etc.) c	ontinuously for the		
☐ Inoperable metastati	c breast cancer	<u>OR</u> □ In	noperable metastatic mamn	nary cano	eer			
•			y for treatment of this cond					
ii. Will the patient found to be pres		• •	a every 6 months and be ac	lvised to	discontinue te	estosterone if		
iii. Will the patient's liver functions tests be monitored every 6 months? □Yes □No								
iv. Will the patient's hematocrit levels be monitored every 6 months? □Yes □No								
	been on testoster xcluding samples		any dosage form (injectio No	n, topica	l, oral, etc.) co	ontinuously for the		

PLEASE PROCEED TO PAGE 2 FOR ADDITIONAL DIAGNOSES

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PAGE 2 - PHYSICIAN COMPLETES					
Patient Name:	DOB:	Patient ID: R			
☐Deficiency of testosterone	<u>OR</u> □Hypogonadisn	n <u>OR</u> □Low testosterone (Low	ow T) OR		
☐Testicular hypofunction	OR □Androgen deficie	ency			
	n testosterone therapy in any ng samples? <i>Please select and</i>	y dosage form (injection, topical, o	ral, etc.) continuously for the		
$\square$ <b>NO</b> – this is <b>INITIA</b>	ATION of therapy, please a	inswer the following questions:			
1) Has the patient h	nad two morning total testos	sterone levels less than 300 ng/dL o	on different days? □Yes □N		
2) What is the patie	ent's hematocrit?	%	ested		
3) Does the patient	have a current diagnosis of	prostate cancer?  \( \subseteq \text{Yes} \) \( \subseteq \text{No} \)			
4) Does the patient	have palpable prostate nod	ules? □Yes □No			
, ·	had a prostatectomy? \(\sigma\)Yes	s □No* state specific antigen (PSA)?	ng/ml DNot tested		
6) Does the patient	have a concurrent diagnosi	s of benign prostate hyperplasia (B worsening symptoms of BPH?	BPH)? □Yes* □No		
,	have a diagnosis of sleep a e patient being treated for th	pnea? □Yes* □No eir sleep apnea? □Yes □No			
	er assessed the patient for the No	heir cardiovascular risk for myocar	dial infarction (MI), angina, or		
10) <b>Natesto Reque</b> anatomy? □Ye		ny chronic nasal conditions or alter	rations in nasal		
$\Box$ <b>YES</b> – this is a PA r	renewal for <b>CONTINUAT</b>	ION of therapy, please answer the	following questions:		
1) Does the patient	have a total testosterone le	vel 800 ng/dL or less? □Yes □	No		
2) Has the patient h	had a prostatectomy?   Yes	s 🗖 No			
· · · · · · · · · · · · · · · · · · ·	•	s of benign prostate hyperplasia (B n BPH worsened since beginning test			
4) Will the patient's	s prostate specific antigen (	PSA) level be tested every 12 mon	ths? □Yes □No		
5) Will the patient's	s serum testosterone concer	ntrations be monitored every 12 mo	onths?  \( \subseteq \text{Yes} \) \( \subseteq \text{No} \)		
6) Will the patient's	s hematocrit levels be mon	itored every 12 months?   Yes	□No		
7) Has the prescribe or stroke? □Yes		or their cardiovascular risk for myo	cardial infarction (MI), angina		
☐ Other (please specify):					

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