SENTARA HEALTH PLANS

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions:</u> The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-844-668-1550</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. <u>If information provided is not complete, correct, or legible, authorization can be delayed</u>.

For Medicare Members: Medicare Coverage for outpatient (Part B) drugs is outlined in the Medicare Benefit Policy Manual (Pub. 100-2), Chapter 15, §50 Drugs and Biologicals. In addition, National Coverage Determination (NCD) and Local Coverage Determinations (LCDs) may exist and compliance with these policies is required where applicable. They can be found at: https://www.cms.gov/medicare-coverage-database/overview-and-quick-search.aspx. Additional indications may be covered at the discretion of the health plan.

Testosterone Replacement Therapy (TRT) Injectables (MUST be purchased by Physician's Office)

Drug Requested: (Check box below that applies)

	NON-I REFERRED				
□ Aveed [®] (testosterone undecanoate) IM injection (J3145)	□ Azmiro (testosterone cypionate) □ Testopel® (testosterone pellets) (11980/S0189)				
MEMBER & PRESCRIBI	R INFORMATION: Authorization may be delayed if incomplete.				
Member Name:					
Member Sentara #:	Date of Birth:				
Prescriber Name:					
Prescriber Signature:	Date:				
Office Contact Name:					
	Fax Number:				
NPI #:					
DRUG INFORMATION:	Authorization may be delayed if incomplete.				
Drug Form/Strength:					
Dosing Schedule:	Length of Therapy:				
Diagnosis:	ICD Code, if applicable:				
Weight (if applicable):	Date weight obtained:				
☐ Standard Review. In checking	this box, the timeframe does not jeopardize the life or health of the memb				

<u>NOTE</u>: For the <u>hypogonadism indication</u>, testosterone drugs <u>CANNOT</u> be used in conjunction with other erectile dysfunction drugs.

the member's ability to regain maximum function and would not subject the member to severe pain.

Maximum Dosage:

- Aveed 3 mL (750 mg) injected intramuscularly, followed by 3 mL (750 mg) injected after 4 weeks, then 3 mL (750 mg) injected every 10 weeks
- Azmiro 50 to 400 mg administered every two to four weeks as an intramuscular injection
- Testopel 75 mg implantable pellet; 6 pellets per 90-day supply (6 billable units every 90 days)
 - o 11980 = CPT1 code for subcutaneous hormone pellet implantation
 - o S0189 = HCPCS code for testosterone pellet, 75 mg

CLINICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

Length of Authorization: 12 months

- ☐ Member must meet **ONE** of the following:
 - ☐ Member has Partial Androgen Insensitivity Syndrome with male gender identity/gender dysphoria or delayed male puberty

OR

- ☐ Member has hypogonadism confirmed by **BOTH** of the following:
 - □ TWO (2) MORNING (6AM to 11AM) testosterone levels obtained on different dates that are below 300 ng/dL or below the lower limit of normal for the reference range (attach lab results with reference ranges from the laboratory for both)

First testosterone level:	

- □ Repeat testosterone or free testosterone level:
- ☐ Member has at least <u>one</u> specific symptom and at least <u>two</u> non-specific symptoms [<u>NOTE</u>: If '<u>decreased spontaneous erections</u>' is the <u>ONLY</u> symptom documented in chart notes, the request will be denied as testosterone replacement is <u>EXCLUDED</u> from coverage for sexual dysfunction]:

Specific Symptoms : (≥1 of the following)		Non-Specific Symptoms: $(\geq 2 \text{ of the following})$	
	Incomplete or delayed sexual development		Decreased energy, motivation, initiative and
	Reduced sexual desire (libido) and activity		self-confidence
	Decreased spontaneous erections*		Depressed mood
	Breast discomfort, gynecomastia		Poor concentration and memory
	Loss of body hair (axillary, facial, and/or		Sleep disturbances, increased sleepiness
	pubic)		Mild anemia (Hgb 10-11 gm/dL)
	Small testes (<5 mL) or shrinking testes		Reduced muscle bulk and strength due to
	Low or zero sperm count		cachexia
	Height loss, low trauma fracture or low		Increased body fat, BMI
	bone mineral density		Diminished physical or work performance
	Hot flushes, sweats		

	AND
	Member has had an inadequate response, contraindication or intolerance to at least a three-month trial with a topical agent e.g., testosterone gel, testosterone patch, testosterone topical solution, testosterone nasal gel (verified by pharmacy paid claims)
	(Please document date/name of drug)
	AND
	Member has had an inadequate response, contraindication or intolerance to at least a three-month trial of a preferred intramuscular testosterone medication e.g., testosterone cypionate, testosterone enanthate or testosterone undecanoate (verified by pharmacy paid claims)
	(Please document date/name of drug)
Med	ication being provided by (check box below that applies):
	Physician's office OR Specialty Pharmacy
review treatm	gent reviews: Practitioner should call Sentara Pre-Authorization Department if they believe a standard would subject the member to adverse health consequences. Sentara's definition of urgent is a lack of ent that could seriously jeopardize the life or health of the member or the member's ability to regain num function.
	*Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. ** vious therapies will be verified through pharmacy paid claims or submitted chart notes. *