STATE OF WISCONSIN

Division of Medicaid Services F-11092 (09/2018) Wis. Admin. Code § DHS 107.10(2)

FORWARDHEALTH PRIOR AUTHORIZATION / PREFERRED DRUG LIST (PA/PDL) FOR GROWTH HORMONE DRUGS

INSTRUCTIONS: Type or print clearly. Before completing this form, read the Prior Authorization/Preferred Drug List (PA/PDL) for Growth Hormone Drugs Instructions, F-11092A. Providers may refer to the Forms page of the ForwardHealth Portal at www.forwardhealth.wi.gov/WIPortal/Content/provider/forms/index.htm.spage for the completion instructions.

Pharmacy providers are required to have a completed Prior Authorization/Preferred Drug List (PA/PDL) for Growth Hormone Drugs form signed by the prescriber before calling the Specialized Transmission Approval Technology-Prior Authorization (STAT-PA) system or submitting a PA request on the Portal or on paper. Providers may call Provider Services at 800-947-9627 with questions.

SECTION I – MEMBER INFORMATION		
Name – Member (Last, First, Middle Initial)		
2. Member ID Number	3. Date of Birth	n – Member
2. Wellber 15 Nullber	J. Date of Billi	I – Member
SECTION II – PRESCRIPTION INFORMATION		
4. Drug Name	5. Drug Strengt	gth
6. Date Prescription Written	7. Refills	
8. Directions for Use		
9. Name – Prescriber		10. National Provider Identifier (NPI) – Prescriber
9. Name – Flescriber		10. National Flovider identifier (NFI) – Flescriber
11. Address – Prescriber (Street, City, State, Zip+4 Code)		
12. Phone Number – Prescriber		
SECTION III – CLINICAL INFORMATION		
13. Diagnosis Code and Description		
Complete the appropriate section of this form:	7	late Caption III A and
 PA requests for growth hormone drugs (except Serostim of PA requests for Serostim: complete Section III B only. 	or Zorbtive): comple	lete Section III A only.
PA requests for Zorbtive: complete Section III C only.		
SECTION III A – CLINICAL INFORMATION FOR GROWTH	HORMONE DRUG	GS (EXCEPT SEROSTIM OR ZORBTIVE)
14. Is the drug requested a preferred growth hormone drug?		☐ Yes ☐ No
If the drug is a non-preferred growth hormone drug, descri	be the reason for t	the request in the space provided.
		Continue



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SECTION III A – CLINICAL INFORMATION FOR GROWTH HORMONE DRUGS (EXCEP (Continued)	T SERO	STIM OF	R ZORB	TIVE)		
15. Is the prescription for the growth hormone drug written by an endocrinologist or through an endocrinology consultation?		Yes		No		
16. Indicate whether or not growth hormone will be used for each of the following congenital conditions.						
1. Noonan syndrome		Yes		No		
2. Prader-Willi syndrome		Yes		No		
3. SHOX gene deficiency disorder		Yes		No		
4. Turner syndrome		Yes		No		
Note: PA requests for medical conditions not listed above are not available through STA	AT-PA.					
17. If growth hormone will not be used for one of the congenital conditions listed in Element being treated in the space provided.	:16, indi	cate the i	medical	condition that is		

Providers are required to include detailed documentation of the medical work-up and testing used to determine the need for growth hormone treatment. Documentation must include the following, when applicable based on the member's age:

- Detailed endocrinology and medical work-up, including medical problem list, current medication list, and medication history
- Height and weight measurements over time plotted on the most clinically appropriate growth chart(s) for age and gender, including growth velocity, growth percentiles, and Z-scores
- Copies of the most recent insulin-like growth factor-1 (IGF-1) and insulin-like growth factor-binding protein 3 (IGF-BP3) lab reports
- Bone age results
- Thyroid-stimulating hormone level
- Nutrition assessment
- · Any other relevant testing, such as advanced imaging of the hypothalamic-pituitary region, if performed

For growth hormone renewal PA requests, providers should include copies of the most recent endocrinology clinic notes, clinically appropriate height and weight growth charts for age and gender, the most current IGF-1 and IGF-BP3 lab testing results, and the most current bone age when applicable based on the member's age.

Continued

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SECTION III A – CLINICAL INFORMATION (Continued)	FOR GROWTH HOR	MONE DRUGS (EX	CEPT SERC	OSTIM OR 2	ZORB	TIVE)
18. Does the member have a recent stimula	ted response growth h	ormone test?		Yes		No
Indicate the type and results of the most	recent stimulated resp	onse growth hormor	ne test.			
1. Arginine Month	Year	Peak response result ng/mL				
2. Clonidine Month	Year	Peak response resul	lt	ng/mL		
3. Glucagon Month	Year	Peak response result ng/mL				
4. Insulin Month	Year	Peak response result ng/mL				
5.	Month	Year	Peak respor	nse result _		ng/mL
6. U Other:	Month	Year	Peak respor	nse result _		ng/mL
7. U Other:	Month	Year	Peak respor	nse result _		ng/mL
of stimulating agent, a copy of the medic and results from each blood sample take					036 16	veis, the time
SECTION III B – CLINICAL INFORMATION				Vas		No
19. Does the member have a diagnosis of A SECTION III C – CLINICAL INFORMATION				Yes		INO
20. Does the member have a diagnosis of si						
on parenteral nutrition?				Yes		No
SECTION IV – AUTHORIZED SIGNATURE						
21. SIGNATURE – Prescriber			22. Date S	igned		
SECTION V – FOR PHARMACY PROVIDE	RS USING STAT-PA					
23. National Drug Code (11 Digits)		24. Days' Supply Requested (Up to 365 Days)				
25. NPI						
26. Date of Service (mm/dd/ccyy) (For STAT the past.)	Γ-PA requests, the date	e of service may be ι	up to 31 days	s in the futu	ire or ι	up to 14 days in
27. Place of Service						
28. Assigned PA Number						

SECTION VI – ADDITIONAL INFORMATION	
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32. Include any additional information in the space below. Additional diagnostic and clinical information explaining the need for the drug requested may also be included here.	