

Gonadotropins

Vantras (implant/injection) J9225, Supprelin LA (implant) J9226, [histrelin acetate] J1675 Prior Authorization Request Medicare Part B Form

Instructions: * Indicates required information – Form may be returned if required information is not provided. Please fax this request to the appropriate fax number listed at the bottom of the page.

	□ Standard Request– (72 Hours)			Urgent Request (standard time frame could place the member's life, health or ability in serious jeopardy)					
	Date Requested								
					Phone		/ Fax		
MEMBER INFORMATION									
*Name: *ID#: *DOB:									
PRESCRIBER INFORMATION									
*Nai	me:		D □ FNP □ DO □ NP □ PA *Phone:						
*Address:				*Fax:					
DISPENSING PROVIDER / ADMINISTRATION INFORMATION									
*Name: Phone:									
			Fax:						
*Address:Fax: PROCEDURE / PRODUCT INFORMATION									
нс	PC Code	Name of Drug	Dos	o (Wt·	kg Ht:	١	Frequency	End Date if	
			003	c (m	kg m	/	Trequency	known	
	olf odmini		rad			fucion			
Self-administered Provider-administered Home Infusion Chart notes attached. Other important information:									
Diagnosis: ICD10: Description:									
□ Provider attests the diagnosis provided is an FDA-Approved indication for this drug									
CLINICAL INFORMATION									
 New Start or Initial Request: (Clinical documentation required for all requests) Provider has reviewed the attached "Criteria for Approval" and attests the member meets ALL required PA criteria. If not, please provide clinical rationale for formulary exception: 									
 Continuation Requests: (Clinical documentation required for all requests) Patient had an <u>adequate response</u> or <u>significant improvement</u> while on this medication. If not, please provide clinical rationale for continuing this medication: 									
ACKNOWLEDGEMENT									
Request By (Signature Required):									

For questions or assistance, please contact Customer Service at 1-877-672-8620, daily, 8am – 8pm (PST) (TTY users should call 1-800-735-2900).



Prior Authorization Group – Gonadotropin PA

Drug Name(s): SUPPRELIN LA VANTAS HISTRELIN ACETATE

Criteria for approval of Prior Authorization Drug:

- 1. Prescribed for an approved FDA diagnosis (as listed below):
- 2. Member does not have any clinically relevant contraindications, or CMS/Plan exclusions, to the requested drug.
- If the member meets all these criteria, they may be approved by the Plan for the requested drug.
- Quantity limits and Tiering will be determined by the Plan.

Exclusion Criteria: N/A

Prescriber Restrictions: N/A

Coverage Duration: Approvals will be for 12 months

FDA Indications:

Supprelin LA, Vantas

- Prostate cancer, Advanced (palliative treatment)
- Central precocious puberty

Off-Label Uses:

N/A

Age Restrictions: Histrelin acetate (Vantas) is not indicated for use in pediatric patients

Other Clinical Considerations:

Pregnancy; may cause fetal harm and spontaneous abortion

Resources:

https://www.micromedexsolutions.com/micromedex2/librarian/CS/AF0468/ND_PR/evidencexpert/ND_P/evidencexpert/ t/DUPLICATIONSHIELDSYNC/91D855/ND_PG/evidencexpert/ND_B/evidencexpert/ND_AppProduct/evidencexpert/ND_T /evidencexpert/PFActionId/evidencexpert.DoIntegratedSearch?SearchTerm=Histrelin%20Acetate&UserSearchTerm=His trelin%20Acetate&SearchFilter=filterNone&navitem=searchGlobal#