

Prior Authorization Review Panel

CHC-MCO Policy Submission

A separate copy of this form must accompany each policy submitted for review. Policies submitted without this form will not be considered for review.

Plan: PA Health & Wellness	Submission Date: 08/01/2024	
Policy Number: PA.CP.PMN.289	Effective Date: 08/2023 Revision Date: 07/2024	
Policy Name: Fezolinetant (Veozah)		
Type of Submission – <u>Check all that apply</u> :		
 □ New Policy ✓ Revised Policy* □ Annual Review - No Revisions □ Statewide PDL - Select this box when submitting policies y when submitting policies for drug classes included on the Statewise Policies for drug classes f		
*All revisions to the policy <u>must</u> be highlighted using track chan	ges throughout the document.	
Please provide any changes or clarifying information for the pol	icy below:	
3Q 2024 annual review: no significant changes; in Appendix B, removed commercially unavailable brand alternatives; references reviewed and updated.		
Name of Authorized Individual (Please type or print):	Signature of Authorized Individual:	
Craig A. Butler, MD MBA	Cray G. B. Cop	

CLINICAL POLICY

Fezolinetant



Clinical Policy: Fezolinetant (Veozah)

Reference Number: PA.CP.PMN.289

Effective Date: 08/2023 Last Review Date: 07/2024

Description

Fezolinetant (Veozah®) is a neurokinin 3 (NK3) receptor antagonist.

FDA Approved Indication(s)

Veozah is indicated for the treatment of moderate to severe vasomotor symptoms due to menopause.

Policy/Criteria

Provider must submit documentation (such as office chart notes, lab results or other clinical information) supporting that member has met all approval criteria.

It is the policy of PA Health & Wellness® that Veozah is **medically necessary** when the following criteria are met:

I. Initial Approval Criteria

- A. Vasomotor Symptoms (must meet all):
- 1. Diagnosis of vasomotor symptoms associated with menopause;
- 2. Age \geq 18 years;
- 3. Failure of hormonal therapy products (not contraceptives, see Appendix B), unless contraindicated or clinically significant adverse effects are experienced;
- 4. Dose does not exceed 45 mg per day (1 tablet per day).

Approval duration: 12 months

B. Other diagnoses/indications

1. Refer to the off-label use policy if diagnosis is NOT specifically listed under section III (Diagnoses/Indications for which coverage is NOT authorized): PA.CP.PMN.53

II. Continued Therapy

- **A. Diagnosis** (must meet all):
 - 1. Currently receiving medication via PA Health & Wellness benefit and documentation supports positive response to therapy or the Continuity of Care policy (PA. PHARM.01) applies;
 - 2. Member is responding positively to therapy as evidenced (e.g., vasomotor symptom reduction);
 - 3. If request is for a dose increase, new dose does not exceed 45 mg per day (1 tablet per day).

Approval duration: 12 months

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B. Other diagnoses/indications (must meet 1 or 2):

1. Currently receiving medication via PA Health & Wellness benefit and documentation supports positive response to therapy or the Continuity of Care policy (PA. PHARM.01) applies.

Approval duration: Duration of request or 12 months (whichever is less); or

2. Refer to the off-label use policy for the relevant line of business if diagnosis is NOT specifically listed under section III (Diagnoses/Indications for which coverage is NOT authorized): PA.CP.PMN.53

III. Diagnoses/Indications for which coverage is NOT authorized:

A. Non-FDA approved indications, which are not addressed in this policy, unless there is sufficient documentation of efficacy and safety according to the off label use policies – PA.CP.PMN.53

IV. Appendices/General Information

Appendix A: Abbreviation/Acronym Key

FDA: Food and Drug Administration NK3: neurokinin 3

Appendix B: Therapeutic Alternatives

This table provides a listing of preferred alternative therapy recommended in the approval criteria. The drugs listed here may not be a formulary agent and may require prior authorization.

Drug Name	Dosing Regimen	Dose Limit/ Maximum Dose		
Estrogen Products				
estradiol (Climara®, Divigel®, Elestrin®,	Varies by formulation	Varies		
Estrace®, EstroGel®,				
Evamist®,				
Menostar®,				
Minivelle®, Vivelle				
Dot®, Delestrogen®)				
estropipate	0.75 mg PO QD; may titrate if needed	6 mg/day		
	(range: 0.75 to 6 mg/day)			
Menest® (esterified estrogens)	0.3 to 1.25 mg PO QD	1.25 mg/day		
Premarin®	0.3 mg PO QD; may titrate if needed	1.25 mg/day		
(conjugated				
estrogens)				
Premphase®,	1 tablet PO QD	1 tablet/day		
Prempro®				
(conjugated estrogens/				
medroxyprogesterone)				

Therapeutic alternatives are listed as Brand name® (generic) when the drug is available by brand name only and generic (Brand name®) when the drug is available by both brand and generic.

CLINICAL POLICY

Fezolinetant



Appendix C: Contraindications/Boxed Warnings

- Contraindication(s): known cirrhosis; severe renal impairment or end-stage renal disease; concomitant use with CYP1A2 inhibitors
- Boxed warning(s): none reported

V. Dosage and Administration

Indication	Dosing Regimen	Maximum Dose
Vasomotor symptoms	1 tablet PO daily	1 tablet/day

VI. Product Availability

Tablet: 45 mg

VII. References

- 1. Veozah Prescribing Information. Northbrook, IL: Astellas Pharma US, Inc; May 2023. Available at: https://www.veozah.com. Accessed May 29, 2024.
- 2. Stuenkel CA, Davis SR, Gompel A, et al. Treatment of the symptoms of the menopause: an Endocrine Society clinical practice guideline. J Clin Endocrinol Metab. 2015; 100(11): 39754011.
- 3. ClinicalTrials.gov. A study of fezolinetant to treat hot flashes in women going through menopause (Daylight). Available at: https://clinicaltrials.gov/ct2/show/NCT05033886. Accessed May 26, 2023.
- Lederman S, Ottery FD, Cano A, et al. Fezolinetant for treatment of moderate-to-severe vasomotor symptoms associated with menopause (SKYLIGHT 1): a phase 3 randomised controlled study. <u>Lancet</u>. 2023 Apr 1;401(10382):1091-1102. doi: 10.1016/S0140-6736(23)00085-5.
- 5. Johnson KA, Martin N, Nappi RE, et al. Efficacy and safety of fezolinetant in moderate to severe vasomotor symptoms associated with menopause: A phase 3 RCT. *J Clin Endocrinol* Metab. 2023 Jul 14;108(8):1981-1997. doi: 10.1210/clinem/dgad058.
- 6. Clinical Pharmacology [database online]. Elsevier; 2024. Available at: https://www.clinicalkey.com/pharmacology/. Accessed May 29, 2024.

Reviews, Revisions, and Approvals	Date
Policy created	07/2023
3Q 2024 annual review: no significant changes; in Appendix B, removed commercially unavailable brand alternatives; references reviewed and updated.	07/2024