

Vilazodone

Adjudication Guideline

Rule Category:
Pharmaceutical

Ref: No:
2023-PH-30

Version Control:
Version No. V1.0

Effective Date:
30/09/2023

Last Update:
30/08/2023

Approved by:
Daman

Responsible:
Medical Standards
& Research

**Related Adjudication
Guidelines:** NA

Table of Contents

1.	Abstract	3
1.1	For Members.....	3
1.2	For Medical Professionals.....	3
2.	Scope	3
3.	Adjudication Policy.....	3
3.1	Eligibility / Coverage Criteria.....	3
3.2	Requirements for Coverage	4
3.3	Non-Coverage.....	5
3.4	Payment and Coding Rules	5
4.	Denial codes	5
5.	Appendices	5
5.1	References	5
5.2	Revision History	5

1. Abstract

1.1 For Members

Vilazodone is a prescription drug and should be used under the supervision of a doctor experienced in diagnosing and treating indicated for the treatment of major depressive disorder (MDD) in adults.

1.2 For Medical Professionals

Vilazodone is a serotonin partial agonist reuptake inhibitor.

2. Scope

The scope of this adjudication rule is to highlight the medical indications, and coverage details for Vilazodone as per the policy terms and conditions of each health insurance plan administered by Daman.

3. Adjudication Policy

3.1 Eligibility / Coverage Criteria

Vilazodone is a serotonin partial agonist reuptake inhibitor indicated for the treatment of:

1. Major Depressive Disorder.

Dosage and Administration:

- The recommended target dosage for Vilazodone is 20 mg to 40 mg orally once daily with food.
- To achieve the target dosage, titrate Vilazodone as follows:
 - Start with an initial dosage of 10 mg once daily with food for 7 days,
 - Then increase to 20 mg once daily with food.
 - The dose may be increased up to 40 mg once daily with food after a minimum of 7 days between dosage increases.

Missed dose:

It should be taken as soon as the patient remembers. If it is almost time for the next dose, the patient should skip the missed dose and take the next dose at the regular time. Two doses should not be taken at the same time.

Dosage forms and strengths available:

- Tablets: 40mg formulation.

Major Depressive Disorder treatment with Vilazodone:

1. Diagnosis of major depressive disorder.
2. Age \geq 18 years.
3. Failure of two of the following for at least 6 weeks
 - SSRI (selective serotonin reuptake inhibitor),
 - SNRI (serotonin norepinephrine reuptake inhibitor),
 - bupropion,
 - mirtazapine.
4. Dose does not exceed 40 mg (1 tablet) per day.

Dose Continuation:

1. Currently receiving medication or has previously met initial approval criteria.
2. is responding positively to therapy.
3. If request is for a dose increase, new dose does not exceed 40 mg (1 tablet) per day.

Special warnings and precautions for use:

Prior to initiating treatment with Vilazodone or another antidepressant, screen patients for a personal or family history of bipolar disorder, mania, or hypomania.

Most common adverse reactions (incidence \geq 5% and at least twice the rate of placebo): diarrhoea, nausea, vomiting, and insomnia.

WARNINGS AND PRECAUTIONS

- Concomitant use of monoamine oxidase inhibitors (MAOIs) or use within 14 days of stopping MAOIs.
- Patients taking, or within 14 days of stopping, monoamine oxidase inhibitors (MAOIs), including MAOIs such as linezolid or intravenous methylene blue, because of an increased risk of serotonin syndrome.

Paediatric Use:

- The safety and effectiveness of Vilazodone have not been established in paediatric patients for the treatment of major depressive disorder (MDD).

3.2 Requirements for Coverage

- Failure to submit, upon request or when requesting a clinical history, indication the need for testing will result in rejection of claim.
- Kindly code the ICD-10 and the CPT codes to the highest level of specificity
- Eligible clinician specialities

Eligible clinical specialities
Psychiatry

3.3 Non-Coverage

- Not covered for visitor plan
- Not covered for policies without Psychiatry benefit
- Age less than 18 years

3.4 Payment and Coding Rules

Kindly apply DOH payment rules and regulations and relevant coding manuals for ICD, Drugs.

4. Denial Codes

Regulator denial codes with description are elaborated for reference. These are specialized codes directed by regulator, that explains the reason of rejection of the service by DAMAN to the providers.

Code	Code Description
CODE-010	Activity/diagnosis inconsistent with clinician specialty
MNEC-004	Service is not clinically indicated based on good clinical practice
MNEC-003	Diagnoses are not covered

5. Appendices

1. [Authorization form link](#)

5.1 References

1. https://www.accessdata.fda.gov/drugsatfda_docs/label/2021/022567s022lbl.pdf
2. <https://www.rcpsych.ac.uk/mental-health/treatments-and-wellbeing/antidepressants>

5.2 Revision History

Date	Change(s)
30/08/2023	Release of V1.0

Disclaimer

By accessing these Daman Adjudication Guidelines, you acknowledge that you have read and understood the terms of use set out in the disclaimer below:
The information contained in this Adjudication Guideline is intended to outline the procedures of adjudication of medical claims as applied by The National Insurance Company – Daman (hereinafter "Daman"). The Adjudication Guideline is not intended to be comprehensive, should not be used as treatment guidelines and should only be used for the purpose of reference or guidance for adjudication procedures and shall not be construed as conclusive. Daman in no way interferes with the treatment of patient and will not bear any responsibility for treatment decisions interpreted through Daman Adjudication Guideline. Treatment of patient is and remains at all times the sole responsibility of the treating Healthcare Provider. This Adjudication Guideline does not grant any rights or impose obligations on Daman. The Adjudication Guideline and all of the information it contains are provided "as is" without warranties of any kind, whether express or implied which are hereby expressly disclaimed.
Under no circumstances will Daman be liable to any person or business entity for any direct, indirect, special, incidental, consequential, or other damages arising out of any use of, access to, or inability to use or access to, or reliance on this Adjudication Guideline including but without limitation to, any loss of profits, business interruption, or loss of programs or information, even if Daman has been specifically advised of the possibility of such damages. Daman also disclaims all liability for any material contained in other websites linked to Daman website.
This Adjudication Guideline is subject to the laws, decrees, circulars and regulations of Abu Dhabi and UAE. Any information provided herein is general and is not intended to replace or supersede any laws or regulations related to the Adjudication Guideline as enforced in the UAE issued by any governmental entity or regulatory authority, or any other written document governing the relationship between Daman and its contracting parties.
This Adjudication Guideline is developed by Daman and is the property of Daman and may not be copied, reproduced, distributed or displayed by any third party without Daman's express written consent. This Adjudication Guideline incorporates the Current Procedural Terminology (CPT®), which is a registered trademark of the American Medical Association ("AMA") and the CPT codes and descriptions belong to the AMA. Daman reserves the right to modify, alter, amend or obsolete the Adjudication Guideline at any time by providing one month prior notice.