

Division: Pharmacy Policy	Subject: Prior Authorization Criteria
Original Development Date: Original Effective Date:	February 13, 2020
Revision Date:	June 16, 2022, February 20, 2023, March 24, 2025

Amlodipine (NORLIQVA® and KATERZIA®) oral solution/suspension

Clinical PA required (preferred): Norliqva® (amlodipine) solution

Non-Preferred: Katerzia® (amlodipine) suspension

LENGTH OF AUTHORIZATION: 6 months

REVIEW CRITERIA:

- Patient must be ≥ 6 years of age.
- Patient has hypertension OR
- Patient has coronary artery disease
 - Chronic stable angina,
 - o Vasospastic angina (Prinzmetal's or Variant Angina)
 - Angiographically documented coronary artery disease (documented by angiography without heart failure or an ejection fraction <40%).
- Trial and failure of preferred calcium channel blockers or rationale why preferred agents cannot be tried (documentation required).
- If request is for Katerzia, patient must also have documented trial and failure of Norliqva solution (documentation required).

CONTINUATION OF THERAPY

- Patient met initial review criteria; AND
- Documentation of improved clinical response; AND
- Patient has not experienced any treatment-restricting adverse effects; AND
- Dosing is appropriate as per labeling or is supported by compendia.

DOSING AND ADMINISTRATION:

• Refer to product labeling at https://www.accessdata.fda.gov/scripts/cder/daf/

