

Batch No: _____

Ingredients	Mfr	Lot #	Expiry Date	Quantity	Measured	Checked
amlODIPINE besylate 5 mg tablets	Pfizer/Sandoz/ Teva			20		
ORA-Blend	Perrigo			q.s.100 mL		

Additional Information:

Equipment:

mortar and pestle glass stirring rod
graduated measure

Procedure:

Follow your Dept. procedures for risk assessment/training/PPE/equipment/facilities/NAPRA level


1. Soak tablets in a small amount of vehicle in a mortar for a minimum of 30 minutes.
2. Levigate to a smooth paste with a pestle. Continue to levigate as vehicle is added in small amounts until a liquid is formed.
3. Transfer liquid contents from mortar to graduate.
4. Use a small amount of vehicle to rinse mortar and add it to graduate.
5. Use vehicle to q.s. to the final volume. Stir well.
6. Transfer to amber bottle and label.

Quality Control:

Expected Product Appearance	Additional Notes
Light peach coloured suspension	

Storage: Refrigerate
Packaging: amber glass bottles
BUD: 30 days

Sample Label:

	amlODIPINE 1 mg/mL Oral Suspension	
	Lot:	BUD:
	Refrigerate	Shake Well

Date Made/Prepared By/Checked By: _____

Reference:

1. Nahata MC, Morosco RS, Hipple TF. Stability of amlodipine besylate in two liquid dosage forms. J Am Pharm Assoc (Wash). 1999 May-Jun;39(3):375-7.

Formulation Reviewed: June, 2024

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