

TELEPHONE: 514-905-5096 FAX: 514-905-5097 technicalservices@medisca.net

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Suggested Formula Enalaprilat 1.25 mg/mL Intravenous Injection (Solution, 10 mL)	FIN	F 008 729
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SUGGESTED FORMULATION

Ingredient Listing	Qty.	Unit	NDC #	Supplier	Lot Number	Expiry Date
Enalaprilat 0.2% Stock Solution †	6.25	mL				
Sodium Chloride, USP	0.074	g				
Benzyl Alcohol (Parenteral Application), NF	0.09	mL				
Sterile Water for Injection, USP	q.s. to 10.0	mL				
Sodium Hydroxide 10% Solution	As required		(
† Enalaprilat 0.2% Stock Solution						
Enalaprilat, USP	TBD	g	d			
Sterile Water for Injection, USP	45.0	mL				
Sterile Water for Injection, USP	q.s. to 50.0	mL				

SPECIAL PREPARATORY CONSIDERATIONS

Ingredient-Specific Information

Light Sensitive (protect from light whenever possible):

Benzyl Alcohol



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SPECIAL PREPARATORY CONSIDERATIONS (CONTINUED) Suggested Preparatory Guidelines ☐ Non-Sterile Preparation Sterile Preparation Processing Error / To account for processing error, pH testing, sterility and endotoxin testing **Testing Considerations:** considerations during preparation, it is suggested to measure an additional 30 to 40% of the required quantities of ingredients. Special Instruction: This formula may contain one or more Active Pharmaceutical Ingredients (APIs) that may be classified as hazardous, please refer & verify the current NIOSH list of Antineoplastic and Other Hazardous Drugs in Healthcare Settings. At this time, General Chapter <800> Hazardous Drugs – Handling in Healthcare Settings is informational and not compendially applicable unless otherwise specified by regulators and enforcement bodies. For information on the scope, intended applicability, and implementation context for USP General Chapter <800>, see: https://www.usp.org/compounding/general-chapter-hazardous-drugs-handlinghealthcare. This formula must be prepared within the appropriate facilities under adequate environmental conditions, following the necessary guidelines and procedures as stated within USP 797 and USP 800 when handling hazardous drugs. Only trained and qualified personnel must prepare this formula. All heat stable, reusable materials and equipment must be sterilized and depyrogenated by dry heat sterilization at 250°C for 2 hours prior to use. Every batch of final product compounded using this procedure must be sterility and endotoxin tested before being dispensed.

All required personal protective equipment (sterile and hazardous if applicable), such as but not limited to, gowns, aprons, sleeves, gloves both inner and outer if applicable, shoe covers, hairnet, head cap, beard cover, eyewear, appropriate face mask, respirator and face shield, etc., where applicable must be worn at all times. In addition, proper personnel cleansing must be done before entering the buffer or clean area.

If applicable, follow all required procedures for hazardous drug handling including but not limited to procurement, transport, storage, preparation, dispensing, administration, clean up (spills) & disposal.

Filter integrity must be validated by performing a filter stress test. If the test demonstrates that the filter might be defective, the solution must be discarded and remade.

If you are a registered 503B facility, please refer to all relevant guidance documents including but not limited to the Code of Federal Regulations (CFR), Guidance for Industry (GFIs) and Compliance Policy Guides (CPGs).

This procedure requires the use of very small quantities of ingredients. All calculations and preparation techniques must be verified before dispensing the final product.



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SUGGESTED PREPARATION (for 10 mL)

Weigh and / or measure the following ingredients when appropriate:

Ingredient Listing	Qty.	Unit	Multiplication factor (*):	Processing Error	Qty. to measure
Enalaprilat 0.2% Stock Solution †	6.25	mL			
Sodium Chloride, USP	0.074	g			
Benzyl Alcohol (Parenteral Application), NF	0.09	mL	©		
Sterile Water for Injection, USP	q.s. to 10.0	mL			
Sodium Hydroxide 10% Solution	As required		1		
		S	2		
† Enalaprilat 0.2% Stock Solution			0		
Enalaprilat, USP	TBD	g			
Sterile Water for Injection, USP	45.0	mL			
Sterile Water for Injection, USP	q.s. to 50.0	mL			

- Takes into account increased batch size conversions and density conversions, if required.
- § Weigh / measure just prior to use.

Preparatory Instruction

IMPORTANT: All preparatory procedures must be performed using proper Aseptic Technique

1. **Equipment sterilization:**

Following the manufacturer's specifications, sterilize and depyrogenate all heat stable, reusable materials and equipment, then return to ambient temperature.



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2. Ingredient quantification:		
A. Determine the potency of Enalaprilat based on the certificate of analysis:		
		100%
MINUS		
Water Content (from certificate of analysis)	-	%
DIVIDED BY		100
EQUALS		
Quantity of water free Enalaprilat, in decimal	-	
MULTIPLIED BY		
Assay on anhydrous basis result (from certificate of analysis)	-	%
DIVIDED BY		100
EQUALS		
i. Potency of Enalaprilat, in decimal	-	
3. Ingredient quantification:		
A. Determine the quantity (in g) of Enalaprilat to make a Enalaprilat 0.2% Stock Sol	lution, batch si	ze (50 mL):
Quantity of Enalaprilat required for 50 mL		0.100 g
DIVIDED BY		
Potency of Enalaprilat, in decimal (Step 2Ai)	-	
EQUALS		
i. Quantity of Enalaprilat needed for 50 mL	-	g



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4. † Enalaprilat 0.2% Stock Solution preparation:

A. Incrementally add the Enalaprilat (amount determined in step 3Ai) to the Sterile Water for Injection (45.0 mL).

Specifications: Continuously mix.

End result: Homogeneous liquid-like solution.

B. Add additional Sterile Water for Injection to the mixture (Step 4A) to fill to the required batch size (50.0 mL).

Specifications: Continuously mix until all solid particles have completely dissolved.

End result: Homogeneous liquid-like solution.

5. **Medium incorporation:**

A. In the given order, sequentially add the following ingredients to the Enalaprilat 0.2% Stock Solution (6.25 mL *plus* processing error adjustments):

-Benzyl Alcohol (Parenteral Application)

-Sodium Chloride

Specifications: Continuously mix.

End result: Homogeneous liquid-like solution.

Note: Add the next ingredient, once the previous one has been completely added and dissolved.

6. Filling to volume:

A. Add additional Sterile Water for Injection to the above mixture to fill to the required batch size (10.0 mL *plus* processing error adjustments).

Specifications: Continuously mix.

End result: Homogeneous liquid-like solution.



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7. pH testing:

- A. Draw an appropriate amount of the mixture (Step 6A).
- B. Test the pH of the sample. It should lie between 6.5 and 7.5.
- C. If the pH < 6.5, carefully add, in a dropwise fashion, the Sodium Hydroxide 10% Solution to the mixture:
 - 1. Draw and transfer 1 or 2 drops of the Sodium Hydroxide 10% Solution to the mixture.
 - 2. Stir for at least 5 minutes to evenly disperse the Sodium Hydroxide 10% Solution.
 - 3. Re-test the pH.
 - 4. Continue to add the Hydrochloric Acid 10% Solution until the pH of 6.5 to 7.5 is obtained.

IMPORTANT: Do not allow the pH to rise above 7.5.

8. Filtering and transferring:

Aseptically filter the solution through a 0.22- μm sterile filter into the recommended dispensing container (see Packaging requirements). Transfer the remainder into a separate dispensing container. This is to be used as the Test sample for sterility and endotoxin testing.

9. Filter integrity test:

Validate filter integrity by performing a filter stress test. If the test demonstrates that the filter might be defective, the solution must be discarded and remade.

10. Terminal Sterilization:

In relation to the chemical composition of the formulation, final packaging, etc., select and validate an end-stage sterilization method and follow the manufacturer's specifications.

11. Sterility and Endotoxin testing:

Validate the Test sample for sterility and endotoxin, in accordance to current USP 797 regulatory guidelines.



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SUGGESTED PRESENTATION

JŪ	GESTED PRI	ESE	NTATION				
	Estimated Beyond-Use Date		24 hours room temperature, 3 days refrigerated, or 45 days frozen, as per USP 797. BUD based on successful endotoxin test result.	Packa Requirem		Sterile, tightly closed, light-resistant unit-dose injection vials.	
		1	Use as directed. Do not exceed dose.	l prescribed	6	Discard in the presence of particulate matter.	
		2	Keep out of reach of children.		7	For intravenous use only.	
	Auxiliary Labels	3	Keep at controlled room temper – 25°C), refrigerated (2°C – 8°C) (-25°C to -10°C).		8	Cap tightly after use.	
		4	Do not use if product changes co	olor.	9	Protect from light.	
		5	Consult your health care practit other prescription or over medications are currently being prescribed for future use.	-the-counter	10		
	Pharmacist Instructions	Add any auxiliary labels specific to the active ingredient to the dispensing container as deemed necessary. Contact your pharmacist in the event of adverse reactions					
	Patient Instructions						



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