

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

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Suggested Formula	Nitrofurantoin 25 mg/5 mL Oral Liquid (Suspension, 100 mL)	FIN	F 001 931v2
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SUGGESTED FORMULATION

Ingredient Listing	Qty.	Unit	NDC #	Supplier	Lot Number	Expiry Date
Nitrofurantoin (Anhydrous), USP	0.500	g				
Stevia Powder	0.10	g				
Glycerin, USP	5.0	mL				
Medisca Oral Suspend (Suspending Vehicle)	50.0	mL				
Medisca Oral Syrup (Flavored Vehicle)	q.s. to 100.0	mL	@)		

SPECIAL PREPARATORY CONSIDERATIONS

Ingredient-Specific Information	4	S				
Hygroscopic (protect from moi	sture whenever possible):	Glycerin, Stevia Powder				
Metal Sensitive (protect from n	netal whenever possible):	Nitrofurantoin				
Light Sensitive (protect from li	ight whenever possible):	Nitrofurantoin				
Suggested Preparatory Guidelines						
Non-Sterile Preparat	ion					
<u>Processing Error /</u> <u>Testing Considerations</u> :		considerations during preparation, it is suggested to the required quantities of ingredients.				
Special Instruction:	Protective apparel, such as a lab of should always be worn.	oat, disposable gloves, eyewear and face-masks				
		f very small quantities of ingredients. All calculations be verified before dispensing the final product.				



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SUGGESTED PREPARATION (for 100 mL)

Weigh and / or measure the following ingredients when appropriate:

Ingredient Listing	Qty.	Unit	Multiplication factor (*):	Processing Error	Qty. to measure
Nitrofurantoin (Anhydrous), USP §	0.500	g			
Stevia Powder §	0.10	g			
Glycerin, USP §	5.0	mL			
Medisca Oral Suspend (Suspending Vehicle)	50.0	mL			
Medisca Oral Syrup (Flavored Vehicle)	q.s. to 100.0	mL	, , , C.		

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

Preparatory Instruction

1. **Powder-liquid preparation:**

- A. Combine and triturate the following ingredients together to form a fine, homogeneous powder blend:
 - -Nitrofurantoin (Anhydrous)
 - -Stevia Powder
- B. Levigate the fine, homogeneous powder blend (Step 1A) with the Glycerin.

End result: Homogeneous liquid-like dispersion.

2. **Medium integration:**

A. Incrementally add the homogeneous liquid-like dispersion (Step 1B) to the Oral Suspend (Suspending Vehicle).

Specifications: Continuously mix.

End result: Homogeneous liquid-like dispersion.

3. **Filling to volume:**

A. Add Oral Syrup (Flavored Vehicle) to the mixture (Step 2A) to fill to the required batch size (100.0 mL *plus* processing error adjustments).

Specifications: Continuously mix.

End result: Homogeneous liquid-like dispersion.



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4. **Product transfer:**

A. Transfer the final product into the specified dispensing container (see "Packaging requirements").

<u>Note</u>: Continuously mix the final product during the transfer process into the recommended dispensing containers in order to maintain homogeneity.

SUGGESTED PRESENTATION

Estimated Beyond-Use Date		14 days, refrigerated, as per USP.	Packaging Requirements		 Tightly closed, light-resistant dispensing bottle. To be administered with a metered dosemeasuring device. 	
	1	Use as directed. Do not exceed dose.	d prescribed	6	Cap tightly after use.	
	2	Keep out of reach of children.		7	Shake well before use.	
Auxiliary Labels	3	Consult your health care practit other prescription or over medications are currently being prescribed for future use.	-the-counter	8	Keep refrigerated. Do not freeze.	
	4	Do not take with alcohol, tranquilizers or other CNS depre		9	Protect from light.	
	5	May impair mental and/or physuse care when operating machinery.				
Pharmacist Instructions	Add any alixiliary labels specific to the active to the dispensing container as deemed necessary					
Patient Instructions	Contact your pharmacist in the event of adverse reactions.					



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4.	Nitrofurantoin (Monograph). <i>United States Pharmacopeia XXVIII / National Formulary 23</i> . Rockville, MD. US Pharmacopeial Convention, Inc. 2004: 1380.
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