

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

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Suggested Formula	Omeprazole 2 mg/mL Oral Liquid (Suspension, 100 mL)	FIN	F 007 304

### **SUGGESTED FORMULATION**

Ingredient Listing	Qty.	Unit	NDC #	Supplier	Lot Number	Expiry Date
Omeprazole (Powder), USP	0.200	g				
Oral Mix Dry Alka, SF (Cherry Flavored)	6.35	g				
Purified Water, USP	60.0	mL				
Purified Water, USP	q.s. to 100.0	mL				





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## SPE

ECIAL PREPARATORY CONSI	DERATIONS	
Ingredient-Specific Information		
Light Sensitive (protect from li	ght whenever possible):	Omeprazole
Moisture Sensitive (protect fro	m humidity whenever possible):	Omeprazole
Oxygen Sensitive (protect from	a oxygen whenever possible):	Omeprazole
Suggested Preparatory Guidelines		- 0
Non-Sterile Preparat	ion Sterile Preparation	C +
<u>Processing Error /</u> <u>Testing Considerations</u> :		considerations during preparation, it is suggested to f the required quantities of ingredients.
Special Instruction:	may be classified as hazardous Antineoplastic and Other Hazar Chapter <800> Hazardous Dru	r more Active Pharmaceutical Ingredients (APIs) that s, please refer & verify the current NIOSH list of rdous Drugs in Healthcare Settings, 2016. General ags – Handling in Healthcare Settings was formally the First Supplement to USP 39-NF 34 and has a date of December 31st, 2019.
	environmental conditions, follow	d within the appropriate facilities under adequate ving the necessary guidelines and procedures as stated when handling hazardous drugs. Only trained and this formula.
	limited to, lab coat, protective	equipment (hazardous if applicable), such as but not sleeves, gloves both inner and outer if applicable, eard cover, eyewear, appropriate face mask, respirator cable must be worn at all times.
		procedures for hazardous drug handling including but sport, storage, preparation, dispensing, administration,
		cility, please refer to all relevant guidance documents a Code of Federal Regulations (CFR), Guidance for Policy Guides (CPGs).
		of 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
Omeprazole (Powder), USP §	0.200	g			
Oral Mix Dry Alka, SF (Cherry Flavored)	6.35	g			
Purified Water, USP	60.0	mL	<b>&amp;</b>		
Purified Water, USP	q.s. to 100.0	mL \			

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

	Preparatory Instruction				
1.	Powder preparation:				
	A. Combine and triturate the following ingredients together to form a fine, homogeneous powder blend:				
	-Omeprazole (Powder) -Oral Mix Dry Alka, SF (Cherry Flavored)				
2.	Powder integration:				
	A. Incrementally add the Purified Water (60.0 mL <i>plus</i> processing error adjustments) to the fine, homogeneous powder blend (Step 1A).				
	Specifications: Continuously mix, using high-shear mixing techniques until all the powder is well dispersed.				
	End result: Homogeneous liquid-like dispersion.				
3.	Filling to volume:				
	A. Allow the suspension to settle for 30-60 seconds and then add additional Purified Water to the mixture (Step 2A) to fill to the required batch size (100.0 mL <i>plus</i> processing error adjustments).				
	Specifications: Continuously mix, using high-shear mixing techniques until the mixture is uniformly suspended.				

# 4. Product transfer:

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

End result: Homogeneous liquid-like dispersion.

Note: Continuously mix the final product during the transfer process in order to maintain homogeneity.



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#### SUGGESTED PRESENTATION 70 days at 4°C, based on - Amber PP bottles. **Packaging** available stability - To be administered with a metered-dose measuring studies Requirements through Medisca\*. device. \*Suggested BUD is based on the exact execution of the indicated ingredient list, quantities and procedures listed within this formulation. Estimated This data is provided for informational purposes only, representing the results of a study of the Note: Beyond-Use Date product stability with various active pharmaceutical ingredients. It does not serve, and may not be construed, as a representation or guarantee of product performance. In all cases the practitioner is advised to consult recognized pharmaceutical compendia and other recognized sources for product formulation and other product characteristics, including stability. MEDISCA Network Inc. makes no warranties or representations with regard to the functioning or appropriateness of this product in any compounded formulation, which use is solely at the discretion and liability of the practitioner. Use as directed. Do not exceed prescribed Protect from light. dose. Consult your health care practitioner if any other May impair mental and or physical ability. prescription or over-the-counter medications are Use care when operating a car or machinery. currently being used or are prescribed for future Auxiliary Labels 8 3 Shake well before use. Keep out of reach of children. Do not take with alcohol, sleep aids, 4 Cap tightly after use. tranquilizers or other CNS depressants. 5 Keep refrigerated. Do not freeze.

# Pharmacist Instructions

Add any auxiliary labels specific to the active to the dispensing container as deemed necessary.

# Patient Instructions

Contact your pharmacist in the event of adverse reactions.

### **REFERENCES**

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- 2. Losec Capsules. In: Canadian Pharmacists Association. *Compendium of Pharmacists and Specialties*, 2015: 1737.
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5. Omeprazole. In: Trissel LA. <i>Trissel's Stability of Compounded Formulations</i> , 5 <sup>th</sup> Edition. American Pharmaceutical Association; 2012: 359.				
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