

7/26/2018; Page 1

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SUGGESTED FORMULATION

Ingredient Listing	Qty.	Unit	NDC #	Supplier	Lot Number	Expiry Date
Omeprazole (Powder), USP 200 mg Bottle	1.00	Bottle				
Oral Mix Dry Alka, SF (Unflavored) 6.35 g Bottle	1.0	Bottle				
Purified Water, USP	5.0	mL				
Purified Water, USP	5.0	mL				
Purified Water, USP	70.0	mL	7			
Purified Water, USP	q.s. to 100.0	mL				

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7/26/2018; Page 1

	Suggested Formula	FIN	F 007 407					
SPE	SPECIAL PREPARATORY CONSIDERATIONS							
	Ingredient-Specific Information							
	Light Sensitive (protect from light whenever possible): Omeprazole							
	Moistu							
	Oxygen	Sensitive (protect from	oxygen whenever possible):	Omeprazole				
	Suggested	Preparatory Guidelines		œ				
		<u>cocessing Error /</u> esting Considerations:	To account for processing error measure an additional 0% of the			it is suggested to		
	Special Instruction: This formula may contain one or more Active Pharmaceutic may be classified as hazardous, please refer & verify th Antineoplastic and Other Hazardous Drugs in Healthcare Chapter <800> Hazardous Drugs – Handling in Healthca published February 1, 2016 in the First Supplement to U delayed official implementation date of December 31 st , 201				curren lettings e Setti P 39-N	nt NIOSH list of s, 2016. General ngs was formally NF 34 and has a		
	This formula must be prepared within the appropriate environmental conditions, following the necessary guidelin within USP 795 and USP 800, when handling hazardou qualified personnel must prepare this formula.				and pro	ocedures as stated		
	All required personal protective equipment (hazardous if applic limited to, lab coat, protective sleeves, gloves both inner and dedicated shoe covers, hairnet, beard cover, eyewear, appropriate and face shield, etc., where applicable must be worn at all times. If applicable, follow all required procedures for hazardous drug I not limited to procurement, transport, storage, preparation, dispe- clean up (spills) & disposal.					ter if applicable,		
	If you are a registered 503B facility, please refer to all relincluding but not limited to the Code of Federal Regular Industry (GFIs) and Compliance Policy Guides (CPGs).							
			s. All calculations al product.					



7/26/2018; Page 2

Suggested Formula	Omeprazole 2 mg/mL Oral Liquid (Suspension, 100 mL)	FIN	F 007 407
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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 200 mg Bottle §	1.00	Bottle			
Oral Mix Dry Alka, SF (Unflavored) 6.35 g Bottle	1.0	Bottle			
Purified Water, USP	5.0	mL	6		
Purified Water, USP	5.0	mL			
Purified Water, USP	70.0	mL	L		
Purified Water, USP	q.s. to 100.0	mL	2		

* Takes into account increased batch size conversions and density conversions, if required.

§ Weigh / measure just prior to use.

Preparatory Instruction

1. **Powder preparation:**

A. Gently tap the Omeprazole (Powder) 200 mg Bottle and transfer the contents into the Oral Mix Dry Alka, SF (Unflavored) 6.35 g Bottle.

Specifications: Gently shake the bottle horizontally until homogeneous.

B. Rinse the Omeprazole (Powder) 200 mg Bottle, including the seal liner with Purified Water (5.0 mL) **TWICE** and transfer into the Oral Mix Dry Alka, SF (Unflavored) 6.35 g Bottle (Step 1A).

2. **Powder integration:**

A. Incrementally add the Purified Water (70.0 mL) into the Oral Mix Dry Alka, SF (Unflavored) 6.35 g Bottle (Step 1B). Close the cap and gently shake the bottle vertically 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).

Specifications: Close the cap and shake vigorously in a vertical motion until the mixture is uniformly suspended.

End result: Homogeneous liquid-like dispersion.



7/26/2018; Page 3

	ggested Formula	Omeprazole 2 mg/mL Oral Liquid (Suspension, 100 mL)	FIN	F 007 407		
4 Product transfor:						

Product transfer:

Use press-in bottle adapter and seal the bottle with the use of the child resistant cap. (see "Packaging Requirements").

SUGGESTED PRESENTATION

		Amber PP bottles: 70 days at 4°C, based on available stability studies through Medisca. To be administered with a metered-dose measuring device.*					
Estimated		*Suggested BUD is based on the <u>exact</u> execution of the indicated ingredient list, quantities and procedures listed within this formulation.					
Beyond-Use I & Packaging Requiremen		<u>Note</u> : This data is provided for informational purposes only, representing the results of a study of the 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.					
	1	Use as directed. Do not exceed prescribed dose.	6	Protect from light.			
Auxiliary Labels	2	May impair mental and or physical ability. Use care when operating a car or machinery.	7	Consult your health care practitioner if any other prescription or over-the-counter medications are currently being used or are prescribed for future use.			
Labels	3	Shake well before use.	8	Keep out of reach of children.			
	4	Do not take with alcohol, sleep aids, tranquilizers or other CNS depressants.	9	Cap tightly after use.			
	5	Keep refrigerated. Do not freeze.					
Pharmacist Instructions	Ado	dd any auxiliary labels specific to the active to the dispensing container as deemed necessary.					
Patient Instructions	Cor						



7/26/2018; Page 4

		Suggested Formula Omeprazole 2 mg/mL Oral Liquid (Suspension, 100 mL)		FIN	F 007 407				
REF	REFERENCES								
		Suspensions. In: Allen, LV, Jr. <i>The Art, Science, and Technology of Pharmaceutical Compounding Fifth Edition.</i> American Pharmacists Association; 2016: 317.							
		Omeprazole. In: Sweetman SC, ed. <i>Martindale: The Complete Drug Reference, 36th Edition</i> . London, England: The Pharmaceutical Press; 2009: 1753.							
		Omeprazole (Monograph). In: O'Neil MJ. <i>The Merck Index 15th Edition</i> . Whitehouse Station, NJ: Merck & Co, Inc.; 2013: Monograph #6939.							
		Omeprazole. In: Trissel LA. <i>Trissel's Stability of Compounded Formulations</i> , 5 th Edition. American Pharmaceutical Association; 2012: 359.							
		 Omeprazole (Monograph). United States Pharmacopeia XL / National Formulary 35. Rockville, MD. US Pharmacopeia Convention, Inc. 2017: 5433. 							
		Omeprazole Systemic. Thomson Micromedex. USP DI – Drug Information for the Health Care Professional, 26 th Edition. Taunton, MA: US Pharmacopeial Convention, Inc; 2006: 2253.							
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