

4/14/2017; Page 1

Suggested	Bacitracin 400 U/mL, Neomycin 3.5 mg/mL, Polymyxin B Sulfate 5000 U/mL	FIN	F 007 031v2
Formula	Topical Wound Liquid (Solution, 500 mL)	1.114	1.007.03172

SUGGESTED FORMULATION

Ingredient Listing	Qty.	Unit	NDC #	Supplier	Lot Number	Expiry Date
Bacitracin (Micronized), USP	200 000	Units				
Neomycin Sulfate, USP	TBD					
Polymyxin B Sulfate, USP	2 500 000	Units				
Sodium Chloride, USP	3.97	g				
Benzalkonium Chloride Solution (50%), NF	0.2	mL	(P)			
Sterile Water for Injection, USP	400.0	mL				
Sterile Water for Injection, USP	q.s. to 500.0	mL				
Sodium Hydroxide 10% Solution	As required					
Hydrochloric Acid 10% Solution	As required	Ś	4			
	ANE ANE		0			



4/14/2017; Page 2

2			Neomycin 3.5 mg/mL, Polymyxin I (Solution, 500 mL)	B Sulfate 5000 U/mL	FIN	F 007 031v2
	IAL PREPARATORY		DERATIONS			
In	gredient-Specific Inform	nation				
	Light sensitive (protect from light whenever possible):Neomycin Sulfate, Polymyxin B Sulfate, Benzalkonium Chloride Solution					
	Hygroscopic (protect f	from mois	sture whenever possible):	Bacitracin, Neomycin Sulfate, Polymyxin B Sulfate Benzalkonium Chloride Solution		
	Air sensitive (protect f	from air v	vhenever possible):	Benzalkonium Chloride Solu	ution	
	Metal reactive (do not	allow to	come into contact):	Benzalkonium Chloride Solu	ution	
	Oxygen Sensitive (pro	tect from	oxygen whenever possible):	Neomycin Sulfate		
<u>Su</u>	uggested Preparatory Gu	<u>idelines</u>		54		
	Non-Sterile	Preparati	on Sterile Preparation	H.		
	Processing Error / Testing Considerations:To account for processing error, pH testing sterility and endotoxin testing considerations during preparation, it is suggested to measure an additional 1 to 3% of the required quantities of ingredients.					
	Special Instruction	<u>on</u> :	This formula must be prepared w environmental conditions, follow within USP 797. Only trained and	ing the necessary guidelines a	nd pro	cedures as stated
			All heat stable, reusable materials by dry heat sterilization at 250°C		ized a	nd depyrogenated
			Every batch of final product com endotoxin tested before being dis		must b	be sterility and
			Protective apparel, such as a steri eyewear and face-masks should a cleansing must be done before en	lways be worn. In addition, p	roper	
			Filter integrity must be validated demonstrates that the filter might remade.			
			This procedure requires the use o and preparation techniques must			



4/14/2017; Page 3

Suggested FormulaBacitracin 400 U/mL, Neomycin 3.5 mg/mL, Polymyxin B Sulfate 5000 U/mL Topical Wound Liquid (Solution, 500 mL)FINF 007	031v2
---	-------

SUGGESTED PREPARATION (for 500 mL)

Weigh and / or measure the following ingredients when appropriate:

Ingredient Listing	Qty.	Unit	Multiplication factor ^(*) :	Processing Error	Qty. to measure
Bacitracin (Micronized), USP §	200 000	Units			
Neomycin Sulfate, USP §	TBD				
Polymyxin B Sulfate, USP §	2 500 000	Units	•		
Sodium Chloride, USP §	3.97	g			
Benzalkonium Chloride Solution (50%), NF §	0.2	mL	マイ		
Sterile Water for Injection, USP §	400.0	mL	8		
Sterile Water for Injection, USP §	q.s. to 500.0	mL	0		
Sodium Hydroxide 10% Solution §	As required	H			
Hydrochloric Acid 10% Solution §	As required				

* 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.



Suggested Formula	Bacitracin 400 U/mL, Neomycin 3.5 mg/mL, Polymyxin B Sulfate 5000 U/mL Topical Wound Liquid (Solution, 500 mL)	FIN	F 007 031v2
2. <u>Ingre</u>	edient quantification:		
A. I	Determine the potency of Bacitracin (Micronized) based on the certificate of analysis:		
		100%	%
Ν	AINUS		
I	Loss on drying (from certificate of analysis)		%
I	DIVIDED BY	100	
E	EQUALS		
	Quantity of dried Bacitracin (Micronized), in decimal		
N	AULTIPLIED BY		
A	Assay on dried basis (from Certificate of Analysis)		units/mg
E	EQUALS		
i	Potency of Bacitracin (Micronized) in (units/mg)		units/mg



ggested Formula	Bacitracin 400 U/mL, Neomycin 3.5 mg/mL, Polymyxin B Sulfate 5000 U/mL Topical Wound Liquid (Solution, 500 mL)	FIN	F 007 031v2
Ingre	edient quantification:		
	Determine the quantity (in g) of Bacitracin required to make a 500 mL batch of Bacitrac Wound Liquid:	in 400	U/mL Topical
Q	Quantity of Bacitracin (Micronized) (in Units) required for 500 mL	200 00	00 U
	DIVIDED BY		
В	Bacitracin (Micronized) potency in (units/mg) (Step 2Ai)	·	U/mg
E	QUALS		
i	Quantity of Bacitracin (Micronized) (in milligrams) needed for 500 mL		mg
Ν	IULTIPLIED BY		
Ν	Aultiplication factor – milligrams to grams	0.0	001
E	QUALS		
ii	Quantity of Bacitracin (Micronized) (in grams) needed for 500 mL		g
Ν	IULTIPLIED BY		
Р	rocessing error adjustments (1 to 3%)	1.01 to	o 1.03
E	QUALS		
ii	i Quantity of Bacitracin (Micronized) needed <i>plus</i> processing error adjustments		g



Suggeste Formul		FIN	F 007 031v2
4. <u>In</u>	redient quantification:		
А.	Determine the potency of Neomycin Sulfate based on the certificate of analysis:		
	MINUS	100	0%
	Loss on drying (from certificate of analysis)		%
	DIVIDED BY	10)
	EQUALS		
	Quantity of dried Neomycin Sulfate, in decimal		
	MULTIPLIED BY		
	Assay (base equivalent) on dried basis result (from certificate of analysis)		μg/mg
	MULTIPLIED BY (Multiplication factor – μg to grams /mg to grams)	0.0	01
	EQUALS		
	i. Potency of Neomycin Sulfate (Base equivalent) in g/g		



0.	gested ormula	Bacitracin 400 U/mL, Neomycin 3.5 mg/mL, Polymyxin B Sulfate 5000 U/mL Topical Wound Liquid (Solution, 500 mL)	FIN	F 007 031v2
5.	Ingr	edient quantification:		
		Determine the quantity (in g) of Neomycin Sulfate required to make a 500 mL batch of N 8.5 mg/mL Topical Wound Liquid:	eomyo	in (Base)
	(Quantity of Neomycin (Base) required for 500 mL		1.750 g
	Ι	DIVIDED BY		
	I	Potency of Neomycin Sulfate (base equivalent), in g/g (Step 4Ai)	-	
	I	EQUALS		
	i	. Quantity of Neomycin Sulfate needed for 500 mL	-	g
	N	MULTIPLIED BY		
	I	Processing error adjustments (1 to 3%)	1	.01 to 1.03
	I	EQUALS		
	i	i. Quantity of Neomycin Sulfate needed <i>plus</i> processing error adjustments	_	g



uggested Formula	Bacitracin 400 U/mL, Neomycin 3.5 mg/mL, Polymyxin B Sulfate 5000 U/mL Topical Wound Liquid (Solution, 500 mL)	FIN	F 007 031v2
	edient quantification: Determine the potency of Polymyxin B Sulfate based on the certificate of analysis:		
N	4INUS	1009	%
E	Coss on drying (from certificate of analysis)	100	%
Ν	Quantity of dried Polymyxin B Sulfate, in decimal MULTIPLIED BY Assay on dried basis (from Certificate of Analysis)		units/mg
E	EQUALS Potency of Polymyxin B Sulfate in (units/mg)		units/mg



Suggested Formula			F 007 031v2		
. Ingr	Ingredient quantification:				
	Determine the quantity (in g) of Polymyxin B Sulfate required to make a 500 mL batch of 5000 U/mL Topical Wound Liquid:	of Poly	myxin B Sulfate		
	Quantity of Polymyxin B Sulfate (in Units) required for 500 mL	2 500	000 U		
I	DIVIDED BY				
I	Polymyxin B Sulfate potency in (units/mg) (Step 6Ai)		U/mg		
F	EQUALS				
i	Quantity of Polymyxin B Sulfate (in milligrams) needed for 500 mL		mg		
ľ	MULTIPLIED BY				
Ν	Multiplication factor – milligrams to grams	0.0	001		
H	EQUALS				
i	i Quantity of Polymyxin B Sulfate (in grams) needed for 500 mL		g		
ľ	MULTIPLIED BY				
I	Processing error adjustments (1 to 3%)	1.01 to	o 1.03		
I	EQUALS				
i	ii Quantity of Polymyxin B Sulfate needed <i>plus</i> processing error adjustments		g		



	ggested ormulaBacitracin 400 U/mL, Neomycin 3.5 mg/mL, Polymyxin B Sulfate 5000 U/mL Topical Wound Liquid (Solution, 500 mL)FINF 007 031v2
8.	Powder-liquid preparation:
	A. In the given order, sequentially add the following ingredients to the Sterile Water for Injection (400.0 mL <i>plus</i> processing error adjustments):
	-Sodium Chloride -Benzalkonium Chloride Solution (50%) -Bacitracin (amount determined in Step 3Aiii) -Neomycin Sulfate (amount determined in Step 5Aii) -Polymyxin B Sulfate (amount determined in Step 7Aiii)
	Specifications: Continuously mix until all solid particles have completely dissolved.
	End result: Homogeneous liquid-like solution.
	Note: Add the next ingredient, once the previous one has been completely added and dissolved.
9.	pH testing:
	A. Draw an appropriate amount of the mixture (Step 8A).
	B. Test the pH of the sample. It should lie between 6.0 and 7.0.
	C. If the pH < 6.0, carefully add, in a dropwise fashion, the Sodium Hydroxide 10% Solution to the mixture:
	 Draw and transfer 1 or 2 drops of the Sodium Hydroxide 10% Solution to the mixture. Stir for at least 5 minutes to evenly disperse the Sodium Hydroxide 10% Solution. Re-test the pH.
	 4. Continue to add the Sodium Hydroxide 10% solution until the pH of 6.0 to 7.0 is obtained.
	IMPORTANT: Do not allow the pH to rise above 7.0.
	D. If the $pH > 7.0$, carefully add, in a dropwise fashion, the Hydrochloric Acid 10% Solution to the mixture:
	 Draw and transfer 1 or 2 drops of the Hydrochloric Acid 10% Solution to the mixture. Stir for at least 5 minutes to evenly disperse the Hydrochloric Acid 10% Solution. Re-test the pH. Continue to add the Hydrochloric Acid 10% Solution until the pH of 6.0 to 7.0 is obtained.
	IMPORTANT: Do not allow the pH to fall below 6.0.



	ggested ormula	Bacitracin 400 U/mL, Neomycin 3.5 mg/mL, Polymyxin B Sulfate 5000 U/mL Topical Wound Liquid (Solution, 500 mL)	FIN	F 007 031v2		
10.	10. Filling to volume:					
	A. Add additional Sterile Water for Injection to the above mixture to fill to the required batch size (500.0 mL <i>plus</i> processing error adjustments).					
	Specifications: Continuously mix.					
	E	and result: Homogeneous liquid-like solution.				
11.	Filte	ring 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.					
12.	Filte	r integrity test:	-			
		ate filter integrity by performing a filter stress test. If the test demonstrates that the filter on must be discarded and remade.	might	be defective, the		
13.	Steri	lity testing:				
	Valid	ate the Test sample for sterility and endotoxins, in accordance to current USP 797 regula	atory g	uidelines.		



4/14/2017; Page 12

Suggested	Bacitracin 400 U/mL, Neomycin 3.5 mg/mL, Polymyxin B Sulfate 5000 U/mL			
Formula	Topical Wound Liquid (Solution, 500 mL)	FIN		

F 007 031v2

GGESTED PRI	ESE	NTATION				
Estimated Beyond-Use Date		7 days, refrigerated, as per USP 797. BUD based on a successful sterility and endotoxin test result.			Sterile, tightly closed, light-resistant dispersin bottle with sterile topical applicators.	
	1	Use as directed. Do not exceed pr dose.	rescribed	7	Keep refrigerated. Do not freeze.	
	2	Keep out of reach of children.		8	Consult your health care practitioner if an prescription or over-the-counter medications ar currently being used or are prescribed for futur use.	
Auxiliary Labels	3	Equilibrate to room temperature before use.		9	Cap tightly after use.	
	4	Do not use if product changes color.		10	Discard in the presence of particulate matter	
	5	For topical use only.		11	Protect from light.	
	6	May impair mental and/or physica Use care when operating a car or ma		12	Do not take with alcohol, sleep aids, tranquilizer or other CNS depressants.	
Pharmacist Instructions	Add any auxiliary labels specific to the active incredient to the dispensing container as deemed necessary					
Patient Instructions						



4/14/2017; Page 13

Suggested	Bacitracin 400 U/mL, Neomycin 3.5 mg/mL, Polymyxin B Sulfate 5000 U/mL	FIN	F 007 031v2
Formula	Topical Wound Liquid (Solution, 500 mL)	1 11 4	1 007 05172

REFERENCES

1.	Parenteral Preparations. In: Allen, LV, Jr. <i>The Art, Science and Technology of Pharmaceutical Compounding Fourth Edition</i> . American Pharmaceutical Association; 2012: 341.
2.	Benzalkonium Chloride. In: Rowe RC. <i>Handbook of Pharmaceutical Excipients, 7th Edition</i> . American Pharmaceutical Association; 2012: 60.
3.	Sodium Chloride. In: Rowe RC. <i>Handbook of Pharmaceutical Excipients</i> , 7 th Edition. American Pharmaceutical Association; 2012: 729.
4.	Bacitracin. In: Sweetman SC, ed. <i>Martindale: The Complete Drug Reference</i> , 36 th Edition. London, England: The Pharmaceutical Press; 2009: 210.
5.	Neomycin Sulfate. In: Sweetman SC, ed. <i>Martindale: The Complete Drug Reference, 36th Edition</i> . London, England: The Pharmaceutical Press; 2009: 305.
6.	Polymyxin B Sulfate. In: Sweetman SC, ed. <i>Martindale: The Complete Drug Reference, 36th Edition</i> . London, England: The Pharmaceutical Press; 2009: 317.
7.	Bacitracin (Monograph). In: O'Neil MJ. <i>The Merck Index 15th Edition</i> . Whitehouse Station, NJ: Merck & Co, Inc.; 2013: Monograph #927.
8.	Neomycin (Monograph). In: O'Neil MJ. <i>The Merck Index</i> 15 th Edition. Whitehouse Station, NJ: Merck & Co, Inc.; 2013: Monograph #6539.
9.	Polymyxin (Monograph). In: O'Neil MJ. <i>The Merck Index 15th Edition</i> . Whitehouse Station, NJ: Merck & Co, Inc.; 2013: Monograph #7693.
10.	Bacitracin. In: Trissel LA. Trissel's Stability of Compounded Formulations, 5 th Edition. American Pharmaceutical Association; 2012: 57.
11.	Neomycin Sulfate. In: Trissel LA. Trissel's Stability of Compounded Formulations, 5 th Edition. American Pharmaceutical Association; 2012: 349.
12.	Polymyxin B Sulfate. In: Trissel LA. <i>Trissel's Stability of Compounded Formulations</i> , 5 th Edition. American Pharmaceutical Association; 2012: 396.
13.	Bacitracin (Monograph). United States Pharmacopeia XXXIX / National Formulary 34. Rockville, MD. US Pharmacopeial Convention, Inc. 2016: 2670.
14.	Neomycin Sulfate (Monograph). United States Pharmacopeia XXXIX / National Formulary 34. Rockville, MD. US Pharmacopeial Convention, Inc. 2016: 5011.
15.	Polymyxin B Sulfate (Monograph). <i>United States Pharmacopeia XXXIX / National Formulary 34</i> . Rockville, MD. US Pharmacopeial Convention, Inc. 2016: 5443.
16.	USP <797>. United States Pharmacopeia XXXIX / National Formulary 34. Rockville, MD. US Pharmacopeial Convention, Inc. 2016: 626.

DISCLAIMER: MEDISCA NETWORK INC., HEREBY REFERRED TO AS 'THE NETWORK', HAS PROVIDED THE FORMULA AND INSTRUCTIONS ABOVE AS A MODEL FOR EDUCATIONAL PURPOSES ONLY ON THE BASIS OF THE RECOGNIZED COMPENDIA AND TEXTS REFERENCED AT THE END OF THIS DOCUMENT. THE NETWORK TAKES NO RESPONSIBILITY FOR THE VALIDITY, SCHEDULING OR ACCURACY OF THIS INFORMATION OR FOR ITS SAFETY OR EFFECTIVENESS, NOR FOR ANY USE THEREOF, WHICH IS AT THE SOLE RISK OF THE LICENSED PHARMACIST. ADJUSTMENTS MAY BE NEEDED TO MEET SPECIFIC PATIENT NEEDS AND IN ACCORDANCE WITH A LICENSED PRESCRIBER'S PRESCRIPTION. THE PHARMACIST MUST EMPLOY APPROPRIATE TESTS TO DETERMINE THE STABILITY OF THIS SUGGESTED FORMULA. THE NETWORK CANNOT BE HELD LIABLE TO ANY PERSON OR ENTITY CONCERNING CLAIMS, LOSS, OR DAMAGE CAUSED BY, OR ALLEGED TO BE CAUSED BY, DIRECTLY OR INDIRECTLY, THE USE OR MISUSE OF THE INFORMATION CONTAINED IN THIS SUGGESTED FORMULA. IN ALL CASES IT IS THE RESPONSIBILITY OF THE LICENSED PHARMACIST TO KNOW THE LAW, TO COMPOUND ANY FINISHED PRODUCT AND TO DISPENSE THESE PRODUCTS IN ACCORDANCE WITH FEDERAL AND STATE LAW.