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Suggested	Fluticasone Propionate 1%, Itraconazole 2% Topical Ointment (Suspension, 30 g)	FIN	F 006 360
Formula	Funcasone Fropionate 1%, firaconazore 2% Topical Ontiment (Suspension, 50 g)	THN	F 000 300

SUGGESTED FORMULATION

Ingredient Listing	Qty.	Unit	NDC #	Supplier	Lot Number	Expiry Date
Fluticasone Propionate, USP	0.300	g				
Itraconazole, USP	0.600	g				
Propylene Glycol, USP	2.4	mL				
Medisca AlpaWash TM	26.61	g				

SPECIAL PREPARATORY CONSIDERATIONS

Ingredient-Specific Information

Light Sensitive (protect from light whenever possible):

Hygroscopic (protect from moisture whenever possible):

Suggested Preparatory Guidelines

Fluticasone Propionate, Itraconazole, Propylene Glycol

Propylene Glycol

Non-Sterile Preparat	ion Sterile Preparation
<u>Processing Error /</u> <u>Testing Considerations</u> :	To account for processing error considerations during preparation, it is suggested to measure an additional 12 to 15% of the required quantities of ingredients.
Special Instruction:	Protective apparel, such as a lab coat, disposable gloves, eyewear and face-masks should always be worn.
	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 Formula	Fluticasone Propionate 1%, Itraconazole 2% Topical Ointment (Suspension, 30 g)	FIN	F 006 360	
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SUGGESTED PREPARATION (for 30 g)

Weigh and / or measure the following ingredients when appropriate:

Ingredient Listing	Qty.	Unit	Multiplication factor ^(*) :	Processing Error	Qty. to measure
Fluticasone Propionate, USP §	0.300	g			
Itraconazole, USP §	0.600	g			
Propylene Glycol, USP §	2.4	mL			
Medisca AlpaWash TM	26.61	g	\odot		

§ 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:
	-Fluticasone Propionate -Itraconazole
	B. Levigate the fine homogeneous powder blend (Step 1A) with the Propylene Glycol.
	End result: Homogeneous liquid-like dispersion.
2.	Powder-liquid to medium integration:
	A. Incrementally add the homogeneous liquid-like dispersion (Step 1B) to the AlpaWash [™] .
	Specifications: Continuously mix, using high-shear mixing techniques.
	End result: Homogeneous gel-like dispersion.
	B. If the final result is gritty, pass it through the ointment mill until it becomes smooth and uniform.
3.	Product transfer:
	Transfer the final product into the specified dispensing container (see "Packaging Requirements").



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Suggested Formula	Flu	ticas	one Propionate 1%, Itraconazole	2% Topical C	intme	ent (Suspension, 30 g)	FIN	F 006 360
GESTED	PRE	ESEI	NTATION					
Est Beyond-Us	tima se D		6 months, as per USP*.	Packa Requirem		 Tightly closed, light-resis To be administered measuring device. 		
		1	Use as directed. Do not exceed dose.	l prescribed	6	Keep in a dry place.		
		2	2 Keep out of reach of children.		7	Cap tightly after use.		
	uxiliary Labels		Consult your health care practit other prescription or over medications are currently being prescribed for future use.	-the-counter	8	For external use only.		
		4	Keep at room temperature (20°C	C − 23°C).	9	Protect from light.		
		5	May impair mental and/or phys Use care when operating machinery.	-	10	Do not take with alcohol, or other CNS depressants.	sleep a	ids, tranquilize
Pharmac	viet	bur pre nec for	te: This non-sterile formulation rned area. If this formulation spared within the appropriate sessary guidelines and procedur mulation make-up and followin ge sterilization is gamma irradia	will be app facilities und es as stated v g the manuf	lied 1 ler ac vithin actur	to an open wound or buildequate environmental con USP <797>. Also, in consi er's specifications, the sug	rned a ndition ideration gested	rea, it must b s, following th on of the overa method of end

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

 Patient
 Contact your pharmacist in the event of adverse reactions.

 Instructions
 The patient is the pharmacist in the event of adverse reactions.

IMPORTANT: The quantity of API administered is directly dependent on the quantity of product applied.

* The BUD is not later than the time remaining until the earliest expiration date of any API or 6 months,

whichever is earlier.

Pharmacist

Instructions

on a successful sterility test result.



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