

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

12/31/2022; Page 1

Suggested Formula	Sildenafil 22 mg Oral Transmucosal Films (Solid Suspension, 30 Films)	FIN	F 009 667

# **SUGGESTED FORMULATION**

Ingredient Listing	Qty.	Unit	NDC #	Supplier	Lot Number	Expiry Date
Sildenafil Citrate, USP**	0.924	g				
Bitterness Reducing Agent (NF-01) (Natural) (Powder)	0.30	g				
Colloidal Silicon Dioxide, NF	0.50	g				
Menthol (Crystals) (Levorotatory) (Natural), USP	0.10	g				
Sucralose, NF	0.075	g				
Purified Water, USP	8.0	mL				
NovaFilm™ Gel Base	18.00	g		+		
Purified Water, USP	q.s. to 30.0	mL	5	_		

<sup>\*\*</sup>Note: Sildenafil Citrate 0.924 g is equivalent to Sildenafil 0.660 g.



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

12/31/2022; Page 2

Suggested Formula Sildenafil 22 mg Oral Transmucosal Films (Solid Suspension, 30 Films) FIN F 009 667

# SPECIAL PREPARATORY CONSIDERATIONS Ingredient-Specific Information *Hygroscopic* (protect from moisture whenever possible): Sildenafil Citrate, Colloidal Silicon Dioxide *Light Sensitive* (protect from light whenever possible): NovaFilm<sup>TM</sup> Gel Base Suggested Preparatory Guidelines Non-Sterile Preparation Sterile Preparation Processing Error / To account for processing error considerations during preparation, it is suggested to **Testing Considerations:** measure an additional 12 to 15% of the required quantities of ingredients. This formula may contain one or more Active Pharmaceutical Ingredients (APIs) that Special Instruction: may be classified as hazardous, please refer & verify the current NIOSH list of Antineoplastic and Other Hazardous Drugs in Healthcare Settings. At this time, General Chapter <800> Hazardous Drugs – Handling in Healthcare Settings is informational and not compendially applicable unless otherwise specified by regulators and enforcement bodies. For information on the scope, intended applicability, and implementation context for USP General Chapter <800>, see: https://www.usp.org/compounding/general-chapter-hazardous-drugs-handlinghealthcare. This formula must be prepared within the appropriate facilities under adequate environmental conditions, following the necessary guidelines and procedures as stated within USP 795 and USP 800, when handling hazardous drugs. Only trained and qualified personnel must prepare this formula. All required personal protective equipment (hazardous if applicable), such as but not limited to, lab coat, protective sleeves, gloves both inner and outer if applicable, dedicated shoe covers, hairnet, beard cover, eyewear, appropriate face mask, respirator and face shield, etc., where applicable must be worn at all times. If applicable, follow all required procedures for hazardous drug handling including but not limited to procurement, transport, storage, preparation, dispensing, administration, clean up (spills) & disposal. If you are a registered 503B facility, please refer to all relevant guidance documents including but not limited to the Code of Federal Regulations (CFR), Guidance for Industry (GFIs) and Compliance Policy Guides (CPGs). This procedure requires the use of very small quantities of ingredients. All calculations

and preparation techniques must be verified before dispensing the final product.



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

12/31/2022; Page 3

Suggested Formula	Sildenafil 22 mg Oral Transmucosal Films (Solid Suspension, 30 Films)	FIN	F 009 667
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# **SUGGESTED PREPARATION (for 30 Films)**

Weigh and / or measure the following ingredients when appropriate:

Ingredient Listing	Qty.	Unit	Multiplication factor (*):	Processing Error	Qty. to measure
Sildenafil Citrate, USP §	0.924	g			
Bitterness Reducing Agent (NF-01) (Natural) (Powder)	0.30	g			
Colloidal Silicon Dioxide, NF §	0.50	g			
Menthol (Crystals) (Levorotatory) (Natural), USP	0.10	g			
Sucralose, NF	0.075	g	7,1		
Purified Water, USP	8.0	mL	2		
NovaFilm™ Gel Base §	18.00	g	· ·		
Purified Water, USP	q.s. to 30.0	mL			

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

1.	Preparatory Step:	

A. Preheat an appropriate convection oven.

Specifications: Temperature: 50°C.

# 2. **Powder-liquid preparation:**

A. By geometric addition, combine and triturate the following ingredients together to form a fine, homogeneous powder blend:

**Preparatory Instruction** 

- -Sildenafil Citrate
- -Bitterness Reducing Agent (NF-01) (Natural) (Powder)
- -Colloidal Silicon Dioxide
- -Menthol (Crystals) (Levorotatory) (Natural)
- -Sucralose
- B. Levigate the fine, homogeneous powder blend (Step 2A) with the Purified Water (8.0 mL *plus* processing error adjustments).

End result: Homogeneous liquid-like dispersion.



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

12/31/2022; Page 4

Formula   Sildenafil 22 mg Oral Transmucosal Films (Solid Suspension, 30 Films)   FIN   F 009 667
---

### 3. **Medium incorporation:**

A. Incrementally add the homogeneous liquid-like dispersion (Step 2B) to the NovaFilm™ Gel Base.

Specifications: Continuously mix, using high-shear mixing techniques.

**NOTE**: Ensure mixing process does not incorporate any air.

End result: Homogeneous gel-like dispersion.

### 4. **Filling to volume:**

A. Add additional Purified Water to the mixture (Step 3A) to fill to the required batch size (30.0 mL *plus* processing error adjustments).

Specifications: Continuously mix, using high-shear mixing techniques.

NOTE: Ensure mixing process does not incorporate any air.

End result: Homogeneous gel-like dispersion.

### 5. **Mold filling and heating:**

A. Fill the 30 blister mold cavities with 1.00 mL of the homogeneous gel-like dispersion (Step 4A) per cavity. Spread the homogeneous gel-like dispersion in the cavity to a uniform thickness.

**NOTE**: Ensure no air bubbles are added to the mold cavities.

B. Heat the filled blister molds to 50°C for 60 to 90 minutes in the preheated convection oven. Do not overheat.

Specifications: Homogeneous solid dispersion.

### 6. Cooling:

- A. Carefully remove the blister mold from the heated oven.
- B. Allow the films to cool for an additional 15- 30 minutes in the blister molds at controlled temperature and relative humidity.

### 7. Validation technique:

- A. Weigh 6 films separately.
- B. The final weight of each film from Step 7A (not including the weight of the blister mold) shall not be less than 90% and not more than 110% of the theoretically calculated weight 0.170 g in accordance to USP guidelines.



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

12/31/2022; Page 5

Suggested Formula	Sildenafil 22 mg Oral Transmucosal Films (Solid Suspension, 30 Films)	FIN	F 009 667
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8. **Product transfer:** 

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

# **SUGGESTED PRESENTATION**

,	GESTED PRI	-OL	MIATION			
	Estima Beyond-Use D		180 days, controlled room temperature or refrigerator, as per USP 795*.	Packa; Requirem		Manually lock blister molds and put into light-resistant resealable foil pouch.
		1	Use as directed. Do not exceed place.	prescribed	6	Do not take with alcohol, sleep aids, tranquilizers or other CNS depressants.
		2	Keep out of reach of children.		7	Keep at controlled room temperature (20°C – 25°C) OR keep refrigerated (2°C – 8°C). Do not freeze.
	Auxiliary Labels	3	Keep in a dry place.		8	Protect from light.
		4	May impair mental and/or physic Use care when operating a car or n		9	Consult your health care practitioner if any other prescription or over-the-counter medications are currently being used or are prescribed for future use.
		5	Discard container after use.			
	Pharmacist Instructions	Add any allylliary labels specific to the API to the dispensing confainer as deemed necessary				
	Patient Instructions	Co	ntact your pharmacist in the event of	f adverse re	action	ns.

<sup>\*</sup> If the API or any other components in the CNSP have an expiration date that is earlier than the assigned BUD, the expiration date supersedes the assigned BUD and must be the assigned shortest date.



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

12/31/2022; Page 6

Suggested Formula	Sildenafil 22 mg Oral Transmucosal Films (Solid Suspension, 30 Films)	FIN	F 009 667
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