

NASAL DECONGESTANT- phenylephrine hydrochloride tablet, coated
Spirit Pharmaceuticals LLC

Nasal Decongestant

Drug Facts

Active Ingredient (in each tablet)

Phenylephrine HCl 10 mg

Purpose

Nasal decongestant

Uses

• temporarily relieves sinus congestion and pressure • temporarily relieves nasal congestion due to the common cold, hay fever or other upper respiratory allergies

Warnings

Do not use

if you are now taking a prescription monoamine oxidase inhibitor (MAOI) (certain drugs for depression, psychiatric, or emotional conditions, or Parkinson's disease), or for 2 weeks after stopping the MAOI drug. If you do not know if your prescription drug contains an MAOI, ask a doctor or pharmacist before taking this product.

Ask a doctor before use if you have

• heart disease • high blood pressure • thyroid disease • diabetes
• trouble urinating due to an enlarged prostate gland

When using this product do not exceed recommended dose

Stop use and ask a doctor if

• nervousness, dizziness, or sleeplessness occur • symptoms do not improve within 7 days or occur with a fever

If pregnant or breast-feeding,

ask a health professional before use.

Keep out of reach of children.

In case of overdose, get medical help or contact a Poison Control Center right away. (1-800-222-1222)

Directions

adults & children 12 years & over	• take 1 tablet every 4 hours
children under 12 years	• do not take more than 6 tablets in 24 hours ask a doctor

Other information

- store between 20-25°C (68-77°F)

Inactive ingredients

Carnauba wax*, colloidal silicon dioxide*, croscarmellose sodium*, D&C yellow#10 aluminum lake*, dicalcium phosphate*, FD&C Blue#1*, FD&C Red #40 , FD&C Yellow#6*, hypromellose, lactose*, magnesium stearate, microcrystalline cellulose, polyethylene glycol, sodium starch glycolate*, starch*, stearic acid*, talc*, titanium dioxide

*contains one or more of these ingredients

Questions or comments?

1-888-333-9792

Distributed by:

Cabinet Health P.B.C.

Principal Display Panel



**COMPOSTABLE
REFILL POUCH**



CABINET:

Nasal Decongestant Refill

COMPARE TO THE ACTIVE INGREDIENT IN:

Sudafed PE® Sinus Congestion*

Phenylephrine HCl 10 mg

Non-Drowsy

RELIEVES:

SINUS AND NASAL CONGESTION · SINUS PRESSURE

60 TABLETS



TO OPEN:

1. POSITION THUMBS INSIDE THE SHORT FLANGE OVER THE LOCKS.
2. GRIP WHILE PIVOTING THUMBS OUTWARD TO OPEN THE LOCKS.

Tamper evident: do not use if pouch is open

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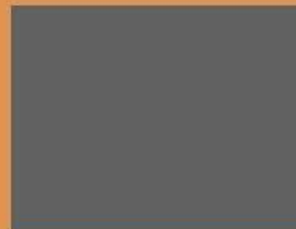
Questions or comments?
 1-888-333-8792

**Get well soon,
 do good now.**

Thanks for helping us on our mission to eliminate single-use plastic from the medicine aisle. By purchasing this compostable refill pouch, you are saving a plastic bottle from a landfill or ocean.

* This product is not manufactured or distributed by McNeil Consumer Healthcare, owner of the registered trademark Sudafed PE® Sinus Congestion

† Less than the limit of detection and consistent with gluten-free diet labeling per FDA



NDC: 68210-4172-1

ACC-1-10011
 Distributed by: Cabinet Health P.B.C.
 Question/Comments? 1-888-333-8792
www.cabinethealth.com

Lot / Expiration ▼



Compostable
 & Plastic Free



Independently
 Quality Tested



Gluten Free
 Verified†

NASAL DECONGESTANT

phenylephrine hydrochloride tablet, coated

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:68210-4234
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PHENYLEPHRINE HYDROCHLORIDE (UNII: 04JA59TNSJ) (PHENYLEPHRINE - UNII:1WS297W6MV)	PHENYLEPHRINE	10 mg

Inactive Ingredients

Ingredient Name	Strength
CARNAUBA WAX (UNII: R12CBM0EIZ)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
CROSCARMELLOSE SODIUM (UNII: M28OL1HH48)	
D&C YELLOW NO. 10 (UNII: 35SW5USQ3G)	
CALCIUM PHENOXIDE (UNII: DRU8G42RVE)	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	
FD&C RED NO. 40 (UNII: WZB9127XOA)	
FD&C YELLOW NO. 6 (UNII: H77VEI93A8)	
HYPROMELLOSES (UNII: 3NXW29V3WO)	
LACTOSE, UNSPECIFIED FORM (UNII: J2B2A4N98G)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)	
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)	
SODIUM STARCH GLYCOLATE TYPE A CORN (UNII: AG9B65PV6B)	
STARCH, POTATO (UNII: 8I089SAH3T)	
STEARIC ACID (UNII: 4ELV7Z65AP)	
TALC (UNII: 7SEV7J4R1U)	
TITANIUM DIHYDRIDE (UNII: 8930U91840)	

Product Characteristics

Color	red	Score	no score
Shape	ROUND	Size	8mm
Flavor		Imprint Code	272;S08;T234
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:68210-	60 in 1 POUCH; Type 0: Not a Combination	11/17/2022	

4234-6	Product	11/17/2022	
Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M012	11/17/2022	

Labeler - Spirit Pharmaceuticals LLC (179621011)

Revised: 12/2024

Spirit Pharmaceuticals LLC