SINUS CONGESTION PE- phenylephrine hcl tablet, film coated WALMART INC.

Equate 44-453

Active ingredient (in each tablet)

Phenylephrine HCl 10 mg

Purpose

Nasal decongestant

Uses

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

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
- diabetes
- thyroid disease
- high blood pressure
- difficulty in urination due to enlargement of the prostate gland

When using this product

do not exceed recommended dosage.

Stop use and ask a doctor if

- nervousness, dizziness, or sleeplessness occur
- symptoms do not improve within 7 days or occur with 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.

Directions

- adults and children 12 years and over: take 1 tablet every 4 hours. Do not take more than 6 tablets in 24 hours.
- children under 12 years: ask a doctor

Other information

- TAMPER EVIDENT: DO NOT USE IF OUTER PACKAGE IS OPENED OR BLISTER IS TORN OR BROKEN
- store at 25°C (77°F); excursions permitted between 15°-30°C (59°-86°F)
- see end flap for expiration date and lot number

Inactive ingredients

croscarmellose sodium, dextrose monohydrate, dibasic calcium phosphate dihydrate, FD&C red #40, lecithin, magnesium stearate, maltodextrin, microcrystalline cellulose, silicon dioxide, sodium carboxymethylcellulose, sodium citrate dihydrate, titanium dioxide

Questions or comments?

1-888-287-1915

Principal display panel

equate™

NDC 79903-076-23

Compare to SUDAFED PE® SINUS CONGESTION active ingredient*

NON-DROWSY Sinus Congestion PE Phenylephrine HCl 10 mg Nasal Decongestant

MAXIMUM STRENGTH

Relieves:

- Sinus Pressure
- Sinus & Nasal Congestion

Actual Size

mg EACH

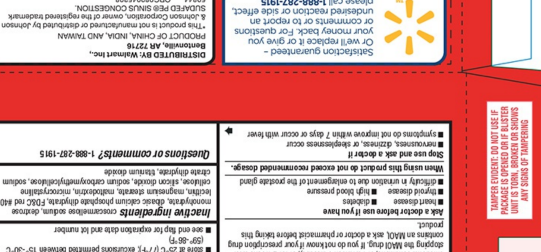
24 TABLETS

TAMPER EVIDENT: DO NOT USE IF PACKAGE IS OPENED OR IF BLISTER UNIT IS TORN, BROKEN OR SHOWS ANY SIGNS OF TAMPERING

Satisfaction guaranteed – Or we'll replace it or give you your money back. For questions or comments please call **1-888-287-1915.**

DISTRIBUTED BY: Walmart Inc., Bentonville, AR 72716

PRODUCT OF CHINA, INDIA, AND TAIWAN *This product is not manufactured or distributed by Johnson & Johnson Corporation, owner of the registered trademark SUDAFED PE® SINUS CONGESTION. 50844 ORG082045308



please call 1-888-287-1915.

When using this product do not exceed recommended dosage.	citrate dihydrate, titanium dioxide		
Ask a doctor before use if you have heart disease dispetes tryroid disease dipy blood pressure tryroid disease dipy blood pressure tryroid dispersion due to enlargement of the prostate gland	Inactive ingredients croscamelloss sodum, dextrose monohydrate, dbasic calcium phosphate dihydrate, FD&C red #40, lecittin, magnesum stearate, maltodextrin, microcrystalline cellulose, slicon dioxide, sodium carboyymethylcellulose, sodium cellulose, slicon dioxide, sodium carboyymethylin, slicon dioxide, sodium carboyymethylin, slicon dioxide, sodium cellulose, slicon dioxide, sodium carboyymethylin, slicon dioxide, sodium cellulose, slicon dioxide, sodium carboyymethylin, sl		
Warnings	Other information		
Do not use if you are now taking a prescription monoamine oxidase	TANPER EVIDENT: DO NOT USE IF OUTER PACKAGE IS OPENED		
inhibitor (MOOI) (certain drugs for deprescion, psychiatric or	OR BLISTER IS TORN OR BROKEN		
emotional conditions, or Parkinson's disease), or for 2 weeks after	a store at 25°C (77°F); excursions permitted between 15°-30°C		
stopping the MOOI drug. If you do not know if your prescription drug	(59°-86°F)		
contains an MAOI, ask a doctor or pharmacist before taking this	= see end flap for expiration date and lot number		
product.	= see end flap for expiration date and lot number		
USeS	Directions		
■ temporarily relieves rasal congestion due to the common cold,	adults and children 12 years and over: take 1 tablet every 4 hours.		
hay fever or other upper respiratory allergies	Do not take more than 6 tablets in 24 hours.		
■ temporarily relieves sinus congestion and pressure	Children under 12 years: ask a doctor		
Purpove (in each tablet) Purpose	If pregnant or breast-feeding, ask a health professional before use.		
Phenylephine HCI 10 mg	Keep out of reach of children. In case of overdose, get medical		
Prenylephine HCI 10 mg	help or contact a Poison Control Center right away.		
Drug Facts	Drug Facts (continued)		

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KEEP OUTER PACKAGE FOR COMPLETE PRODUCT INFORMATION

NDC 79903-076-24

Compare to SUDAFED PE® SINUS CONGESTION gredien

10 mg

EACH

24 TABLETS No Print/ No Varnish Lot #/ Exp. Date



Sinus Congestion PE

Phenylephrine HCl 10 mg **Nasal Decongestant**

MAXIMUM STRENGTH

Relieves: Sinus Pressure Sinus & Nasal Congestion

Equate 44-453

SINUS CONGESTION PE									
phenylephrine hcl tablet, film coated									
Ρ	roduct Info	rmation							
Pi	roduct Type		HUMAN OTC DR	RUG	ltem C	ode (S	ource)	NDC:799	03-076
		istration	ORAL			• • • •	· · · · ·		
	Route of Administration ORAL								
A	ctive Ingred	ient/Active	e Moiety						
		Ingro	edient Name				Basis of S	trength	Strength
	IENYLEPHRINE III:1WS297W6MV)	HYDROCHLOF	RIDE (UNII: 04JA59	TNSJ) (PHEN	NYLEPHR	INE -	PHENYLEPHRINE HYDROCHLORID		10 mg
In	active Ingre	edients							
			Ingredie	nt Name					Strength
CF	ROSCARMELLOS	SE SODIUM (U	NII: M28OL1HH48)						Subigui
	XTROSE MONO								
DI	BASIC CALCIUM	1 PHOSPHATE	DIHYDRATE (UN	III: 07TSZ97	7GEP)				
FC	&C RED NO. 40) (UNII: WZB91	27XOA)						
LE	CITHIN, SOYBE	AN (UNII: 1DI5	6QDM62)						
M	AGNESIUM STE	ARATE (UNII: 7	0097M6I30)						
M		UNII: 7CVR7L4A	2D)						
М	CROCRYSTALLI	NE CELLULOS	5E (UNII: OP1R32E	D61U)					
SI	LICON DIOXIDE	(UNII: ETJ7Z6X	(BU4)						
CA	RBOXYMETHYI	CELLULOSE S	SODIUM, UNSPE	CIFIED (UN	II: K6790	DBS311)		
TR		ATE DIHYDRA	TE (UNII: B22547B	95K)					
тľ	TANIUM DIOXID	E (UNII: 15FIX9	9V2JP)						
_									
	roduct Char			-					
	olor	ree		Score				no score	
	nape	RC	DUND	Size				7mm	
	avor			Imprint C	ode			44;453	
Сс	ontains								
Pa	ackaging								
#	Item Code	P	ackage Descr	iption		Mar	keting Start Date		eting End Date
1	NDC:79903- 076-23	3 in 1 CARTO	N			08/31/2021			
1	NDC:79903- 076-43	24 in 1 BLIST Product	n 1 BLISTER PACK; Type 0: Not a Combination luct		oination				
2	NDC:79903- 076-24	1 in 1 CARTO				08/31/2	2021		
2	NDC:79903- 076-43	24 in 1 BLIST Product	ER PACK; Type 0:	Not a Comb	oination				

Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
OTC Monograph Drug	M012	08/31/2021				

Labeler - WALMART INC. (051957769)

Establishment						
Name	Address	ID/FEI	Business Operations			
LNK International, Inc.		832867837	manufacture(79903-076) , pack(79903-076)			

Establishment

Name	Address	ID/FEI	Business Operations
LNK International, Inc.		832867894	manufacture(79903-076)

Establishment					
Name	Address	ID/FEI	Business Operations		
LNK International, Inc.		868734088	manufacture(79903-076)		

Establishment					
Name	Address	ID/FEI	Business Operations		
LNK International, Inc.		117025878	manufacture(79903-076)		

Revised: 7/2023

WALMART INC.