SUDAFED PE SINUS CONGESTION- diphenhydramine hydrochloride and phenylephrine hydrochloride Johnson & Johnson Consumer Inc.

SUDAFED PE SINUS CONGESTION SUDAFED PE ® DAYTIME

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 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

- store between 20-25°C (68-77°F)
- do not use if blister unit is torn or broken

Inactive ingredients

carnauba wax, D&C yellow no. 10 aluminum lake, FD&C red no. 40 aluminum lake, FD&C yellow no. 6 aluminum lake, magnesium stearate, microcrystalline cellulose, modified starch, polyethylene glycol, polyvinyl alcohol, powdered cellulose, pregelatinized starch, sodium starch glycolate, talc, titanium dioxide

Questions or comments?

call **1-888-217-2117** (toll-free) or **215-273-8755** (collect)

SUDAFED PE ® NIGHTTIME

Drug Facts

Active ingredients (in each tablet)	Purpose
Diphenhydramine HCl 25 mg	Antihistamine
Phenylephrine HCl 10 mg	Nasal decongestant

Uses

- temporarily relieves these symptoms due to hay fever or other upper respiratory allergies:
 - runny nose
 - sneezing
 - itchy, watery eyes
 - itching of the nose or throat
 - nasal congestion
- temporarily relieves these symptoms due to the common cold:
 - runny nose
 - sneezing
 - nasal congestion

• temporarily relieves sinus congestion and pressure

Warnings

Do not use

- to make a child sleepy
- with any other product containing diphenhydramine, even one used on skin
- 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
- a breathing problem such as emphysema or chronic bronchitis
- glaucoma

Ask a doctor or pharmacist before use if you are taking sedatives or tranquilizers

When using this product

- do not exceed recommended dose
- marked drowsiness may occur
- avoid alcoholic drinks
- alcohol, sedatives, and tranquilizers may increase drowsiness
- be careful when driving a motor vehicle or operating machinery
- excitability may occur, especially in children

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 and children 12	 take 1 tablet every 4 hours do not take more than 6
years and over	tablets in 24 hours
children under 12	ack a doctor

Other information

- store between 20-25°C (68-77°F)
- do not use if blister unit is torn or broken

Inactive ingredients

carnauba wax, FD&C blue no. 1 aluminum lake, magnesium stearate, microcrystalline cellulose, modified starch, polyethylene glycol, polyvinyl alcohol, powdered cellulose, pregelatinized starch, sodium starch glycolate, talc, titanium dioxide

Questions or comments?

call **1-888-217-2117** (toll-free) or **215-273-8755** (collect)

PRINCIPAL DISPLAY PANEL

PREVIOUSLY SUDAFED PE ® DAY + NIGHT SINUS CONGESTION NDC 50580-239-01

SUDAFED PE®

SINUS CONGESTION

Phenylephrine HCl Nasal Decongestant

DAYTIME

 NASAL CONGESTION

actual size

12 TABLETS 10 mg each

Diphenhydramine HCl, Phenylephrine HCl Antihistamine, Nasal Decongestant

NIGHTTIME

- NASAL CONGESTION
- RUNNY NOSE

actual size

8 TABLETS | TOTAL: 20 TABLETS

The makers of the SUDAFED® family of products are dedicated to helping you and your family reduce head congestion and sinus pressure.

PREVIOUSLY SUDAFED PE® DAY + NIGHT SINUS CONGESTION NDC 50580-239-01

SUDAFE

SINUS CONGESTION

Phenylephrine HCI Nasal Decongestant

Diphenhydramine HCI, Phenylephrine HCI Antihistamine, Nasal Decongestant

DAYTIME





Does not contain Pseudoephedrine

 NASAI CONGESTION



- NASAL CONGESTION
- RUNNY NOSE

12 TABLETS 10 mg each 8 TABLETS | TOTAL: 20 TABLETS



Distributed by: JOHNSON & JOHNSON CONSUMER INC. McNeil Consumer Healthcare Division

Fort Washington, PA 19034 USA @J&JCI 2018 Daytime: Active ingredient made in Germany Nighttime: Made in Italy

DO NOT USE IF BLISTER UNIT IS TORN OR BROKEN

call 1-888-217-2117 (foll-free) or 215-275-8755 (collect) Questions of continents? 00000

- do not use if blister unit is tom or broken ■ store between 20-25°C (68-77°F)
 - Other information

hildren under	ask adoctor	
dults and children 2 years and over	■ take 1 tablet every 4 hours ■ do not take more tran 6 tablets in 24 hours	
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Directions

or contact a Poison Cortrol Center right away. (1-800-222-1222) Keep out of resch of children. In case of overcose, get medical help If pre gnant or breast-feeding, ask a health professional before use.

Drug Facts (continued)

TOTAL: 20 TABLETS STABLETS

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3. NIGHTTIME

Masal Decongestant

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JO mg each

I Z TABLETS



DAYTIME

Masal Decongestant

Phenylephrine HCI

Olphenhydramine HCl Phenylephrine HCl Antihistamine,

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NDC 20280-528-01

cs|| 1-888-713-7113 (toll-tiee) or 215-235-8355 (collect) Questions or comments?

starch glycolate, talc, thanum dioxide dýcor, pojvviný akohol, powdered cellulose, pregelatinízed starch, sodium magnesium stearate, microcrystalline cellulose, modified starch, polyethylene lake, FD&C red no. 40 aluminum lake, FD&C yellow no. 6 aluminum lake,

- ■symptoms do notimprove within 7 days or occur with a fever ■ nervousness, dzziness, or sleeplessness occur
 - Stop use and ask a doctor if
- excusplity may occur, especially in children
- pe cereţni wueu qiiying a motor vehide or operating machinery. ■ alcohol, sedatives, and tranquilizers may increase drowsiness





SUDAFED PE SINUS CONGESTION

diphenhydramine hydrochloride and phenylephrine hydrochloride kit

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:50580-239

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:50580-239-01	1 in 1 PACKAGE	06/17/2019		

Quantity of Parts

Part # Package Quantity Total Product Quantity

		-
Part 1	2 BLISTER PACK	12
Part 2	2 BLISTER PACK	8

Part 1 of 2

SUDAFED PE SIINUS CONGESTION

phenylephrine hydrochloride tablet, film coated

Product Information

Route of Administration ORAL

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

Inactive Ingredients			
Ingredient Name	Strength		
CARNAUBA WAX (UNII: R12CBM0EIZ)			
D&C YELLOW NO. 10 ALUMINUM LAKE (UNII: CQ3XH3DET6)			
FD&C RED NO. 40 (UNII: WZB9127XOA)			
FD&C YELLOW NO. 6 (UNII: H77VEI93A8)			
ALUMINUM OXIDE (UNII: LMI26O6933)			
MAGNESIUM STEARATE (UNII: 70097M6I30)			
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)			
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)			
POLYVINYL ALCOHOL, UNSPECIFIED (UNII: 532B59J990)			
POWDERED CELLULOSE (UNII: SMD1X3XO9M)			
SODIUM STARCH GLYCOLATE TYPE A (UNII: H8AV0SQX4D)			
TALC (UNII: 7SEV7J4R1U)			
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)			

Product Characteristics			
Color	red	Score	no score
Shape	ROUND	Size	7mm
Flavor		Imprint Code	WL;80;PE
Contains			

Packaging				
#	ltem Code	Package Description	Marketing Start Date	Marketing End Date
1		2 in 1 CARTON		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M012	06/17/2019	

Part 2 of 2

SUDAFED PE SINUS CONGESTION PLUS ALLERGY

diphenhydramine hydrochloride and phenylephrine hydrochloride tablet, film coated

Product Information

Route of Administration ORAL

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
DIPHENHYDRAMINE HYDROCHLORIDE (UNII: TC2D6JAD40) (DIPHENHYDRAMINE - UNII:8GTS82S83M)	DIPHENHYDRAMINE HYDROCHLORIDE	25 mg	
PHENYLEPHRINE HYDROCHLORIDE (UNII: 04JA59TNSJ) (PHENYLEPHRINE - UNII: 1W5297W6MV)	PHENYLEPHRINE HYDROCHLORIDE	10 mg	

Inactive Ingredients		
Ingredient Name	Strength	
CARNAUBA WAX (UNII: R12CBM0EIZ)		
FD&C BLUE NO. 1 ALUMINUM LAKE (UNII: J9EQA3S2JM)		
MAGNESIUM STEARATE (UNII: 70097M6I30)		
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)		
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)		
POLYVINYL ALCOHOL, UNSPECIFIED (UNII: 532B59J990)		
POWDERED CELLULOSE (UNII: SMD1X3XO9M)		
SODIUM STARCH GLYCOLATE TYPE A (UNII: H8AV0SQX4D)		
TALC (UNII: 7SEV7J4R1U)		
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)		

Product Characteristics			
Color	blue	Score	no score
Shape	OVAL	Size	13mm
Flavor		Imprint Code	PE;WL95
Contains			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1		2 in 1 CARTON		
1		4 in 1 BLISTER PACK; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M012	06/17/2019	

Marketing In	Marketing Information		
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M012	06/17/2019	

Labeler - Johnson & Johnson Consumer Inc. (878046358)

Revised: 1/2024 Johnson & Johnson Consumer Inc.