NIGHT TIME SEVERE COLD AND FLU RELIEF- acetaminophen, phenylephrine hydrochloride, doxylamine succinate, dextromethorphan hydrobromide capsule, liquid filled PuraCap Pharmaceutical LLC

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Night Time Severe Cold and Flu Relief

Drug Facts

Active ingredient (in each softgel)

Acetaminophen 325 mg

Dextromethorphan HBr 10 mg

Doxylamine succinate 6.25 mg

Phenylephrine HCl 5 mg

Purposes

Pain reliever/fever reducer

Cough suppressant

Antihistamine

Nasal decongestant

Uses

temporarily relieves common cold/flu symptoms:

- nasal congestion
- sinus congestion & pressure
- cough due to minor throat & bronchial irritation
- cough to help you sleep
- minor aches & pains
- headache
- fever
- sore throat
- runny nose & sneezing
- reduces swelling of nasal passages
- temporarily restores freer breathing through the nose
- promotes nasal and/or sinus drainage

Warnings

Liver warning This product contains acetaminophen. Severe liver damage may occur if you take

- more than 4 doses in 24 hrs, which is the maximum daily amount for this product
- with other drugs containing acetaminophen
- 3 or more alcoholic drinks every day while using this product

Sore throat warning

If sore throat is severe, persists for more than 2 days, is accompanied or followed by fever, headache, rash, nausea, or vomiting, consult a doctor promptly.

Allergy alert: Acetaminophen may cause severe skin reactions. Symptoms may include:

- skin reddening
- blisters
- rash

If a skin reaction occurs, stop use and seek medical help right away.

Do not use

- with any other drug containing acetaminophen (prescription or nonprescription). If you are not sure whether a drug contains acetaminophen, ask a doctor or pharmacist.
- 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.
- to make a child sleep

Ask a doctor before use if you have

- liver disease
- heart disease
- high blood pressure
- thyroid disease
- diabetes
- glaucoma
- cough that occurs with too much phlegm (mucus)
- a breathing problem or chronic cough that lasts or as occurs with smoking, asthma, chronic bronchitis, or emphysema
- trouble urinating due to enlarged prostate gland

Ask a doctor or pharmacist before use if you are

- taking sedatives or tranquilizers
- taking the blood thinning drug warfarin

When using this product

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

Stop use and ask a doctor if

- you get nervous, dizzy or sleepless
- pain, nasal congestion, or cough gets worse or lasts more than 7 days
- fever gets worse or lasts more than 3 days
- redness or swelling is present
- new symptoms occur
- cough comes back or occurs with rash or headache that lasts. These could be signs of a serious condition.

If pregnant or breast-feeding, ask a health professional before use.

Keep out of reach of children.

Overdose warning

Taking more than directed can cause serious health problems. In case of overdose, get medical help or contact a Poison Control Center right away. Quick medical attention is critical for adults as well as for children even if you do not notice any signs or symptoms.

Directions

- take only as directed see **Overdose warning**
- do not exceed 4 doses per 24 hrs

adults & children 12 yrs and over	2 softgels with water every 4 hrs
children 4 to under 12 yrs	ask a doctor
children under 4 yrs	do not use

• when using other Nighttime or Daytime products, carefully read each label to ensure correct dosing

Other information

• store at room temperature and avoid excessive heat

Inactive ingredients

D&C yellow #10, FD&C blue #1, gelatin, glycerin, polyethylene glycol, povidone, propylene glycol, purified water, sorbitol special, and white edible ink

Questions or comments?

Call toll free: **1-888-309-9030**

PRINCIPAL DISPLAY PANEL - Carton Label

Night Time Severe Cold & Flu Relief 24 Softgels

Acetaminophen 325mg, Phenylephrine hydrochloride 5mg, Doxylamine succinate 6.25mg, Dextromethorphan hydrobromide 10mg

NDC 51013-175-04

*Compare to the active ingredients in Vicks® NyQuil® Severe Cold and Flu



NIGHT TIME SEVERE COLD AND FLU RELIEF

acetaminophen, phenylephrine hydrochloride, doxylamine succinate, dextromethorphan hydrobromide capsule, liquid filled

Product Information					
Product T ype	HUMAN OTC DRUG	Item Code (Source)	NDC:51013-175		
Route of Administration	ORAL				
Active Ingredient/Active Moiety					

	Ingredient Name		Basis of Str	ength	Strength	
ACETAMINOPHEN (UN	III: 36209ITL9D) (ACETAMINOPHEN -	: 36209ITL9D) (ACETAMINOPHEN - UNII:36209ITL9D)			325 mg	
	METHO RPHAN HYDRO BRO MIDE (UNII: 9 D2RTI9 KYH)DEXTROMETHORPHANMETHO RPHAN - UNII:7355X3ROTS)HYDRO BRO MIDE				10 mg	
DO XYLAMINE SUCCIN	NATE (UNII: V9BI9B5YI2) (DOXYLAMI	INE - UNII:95QB77JKI	PL) DOXYLAMINE SUC	CINATE	6.25 mg	
PHENYLEPHRINE HYD UNII:1WS297W6MV)	ROCHLORIDE (UNII: 04JA59TNSJ) (I	PHENYLEPHRINE		5 mg		
Inactive Ingredier	its					
	Ingredient Na	ame		St	rength	
D&C YELLOW NO. 10	(UNII: 35SW5USQ3G)					
FD&C BLUE NO. 1 (UN	II: H3R47K3TBD)					
GELATIN (UNII: 2G86Q	N327L)					
GLYCERIN (UNII: PDC6	A3C0OX)					
POLYETHYLENE GLY	COL, UNSPECIFIED (UNII: 3WJQ0SD	W1A)				
POVIDONE (UNII: FZ98	,					
PROPYLENE GLYCOL	(UNII: 6DC9Q167V3)					
WATER (UNII: 059QF0F	KOOR)					
SORBITOL (UNII: 5067	(60A25R)					
Product Character	ristics					
Color	green (clear)	reen (clear) Score		no score		
Shape	capsule (oblong)	Size		20 mm		
Flavor		Imprint Cod	Imprint Code PC		PC22	
Contains						
Packaging						
# Item Code	Package Description Mar		farketing Start Date	Marketing	, End Date	
1 NDC:51013-175-04 2	1 CARTON 06/0		6/09/2016			
1 12	in 1 BLISTER PACK; Type 0: Not a Co	mbination Product				
Marketing Info	rmation					
Marketing Category	Application Number or Monog	raph Citation M	arketing Start Date	Marketing	End Date	
OTC monograph final	part341	06	/09/2016			

Labeler - PuraCap Pharmaceutical LLC (962106329)

Name	Address	ID/FEI	Business Operations
Humanwell PuraCap Pharmaceutical (Wuhan) Co., Ltd		421293287	manufacture(51013-175) , analysis(51013-175)