### SEVERE COLD AND FLU- acetaminophen, dextromethorphan hbr, diphenhydramine hcl, guaifenesin, phenylephrine hcl Wal-Mart Stores Inc

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### Equate 44-042011

### Active ingredients (in each 20 mL) (Daytime)

Acetaminophen 650 mg Dextromethorphan HBr 20 mg Guaifenesin 400 mg Phenylephrine HCl 10 mg

### Purpose

Pain reliever/fever reducer Cough suppressant Expectorant Nasal decongestant

### Active ingredients (in each 20 mL) (Nighttime)

Acetaminophen 650 mg Diphenhydramine HCl 25 mg Phenylephrine HCl 10 mg

### Purpose

Pain reliever/fever reducer Antihistamine/cough suppressant Nasal decongestant

### Uses

- temporarily relieves these common cold and flu symptoms:
  - cough
  - sore throat
  - headache
  - minor aches and pains
  - nasal congestion
  - runny nose and sneezing (Nighttime only)
- temporarily reduces fever
- helps loosen phlegm (mucus) and thin bronchial secretions to rid the bronchial passageways of bothersome mucus and make coughs more productive (Daytime only)
- controls cough to help you get to sleep (Nighttime only)

# Warnings

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

- with other drugs containing acetaminophen
- more than 4,000 mg of acetaminophen in 24 hours
- 3 or more alcoholic drinks every day while using this product

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

- blisters
- rash
- skin reddening

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

**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.

## 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.
- 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 have ever had an allergic reaction to this product or any of its ingredients
- with any other product containing diphenhydramine, even one used on skin (Nighttime only)

## Ask a doctor before use if you have

- heart disease
- diabetes
- thyroid disease
- liver disease
- high blood pressure
- persistent or chronic cough such as occurs with smoking, asthma, chronic bronchitis, or emphysema
- difficulty in urination due to enlargement of the prostate gland
- cough that occurs with too much phlegm (mucus)
- a breathing problem such as emphysema or chronic bronchitis (*Nighttime only*)
- glaucoma (Nighttime only)

# Ask a doctor or pharmacist before use if you are

- taking the blood thinning drug warfarin
- taking sedatives or tranquilizers (Nighttime only)

## When using this product

- do not exceed recommended dosage
- excitability may occur, especially in children (Nighttime only)
- marked drowsiness may occur (Nighttime only)
- avoid alcoholic beverages (Nighttime only)
- alcohol, sedatives, and tranquilizers may increase drowsiness (Nighttime only)
- use caution when driving a motor vehicle or operating machinery (Nighttime only)

## Stop use and ask a doctor if

- nervousness, dizziness, or sleeplessness occur
- 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.

In case of accidental overdose, get medical help or contact a Poison Control Center right away. Prompt medical attention is critical for adults as well as for children even if you do not notice any signs or symptoms.

# If taking NIGHTTIME and DAYTIME products, carefully read each section to ensure correct dosing. Do not take DAY & NIGHT at the same time.

# Directions

- do not take more than directed
- do not take more than 6 doses in any 24-hour period
- mL = milliliter; FL OZ = fluid ounce
- use only enclosed dosing cup designed for use with this product. Do not use any other dosing device.
- adults and children 12 years and over: 20 mL in dosing cup provided every 4 hours
- children under 12 years: do not use

# Other information

- each 20 mL contains: sodium 10 mg (Daytime only), sodium 9 mg (Nighttime only)
- store at 25°C (77°F); excursions permitted between 15°-30°C (59°-86°F)
- use by expiration date on package

# Inactive ingredients (Daytime only)

anhydrous citric acid, disodium edetate, FD&C blue #1, FD&C red #40, flavors, glycerin,

polyethylene glycol, propylene glycol, purified water, sodium benzoate, sodium citrate dihydrate, sodium metabisulfite, sorbitol, sucralose, xanthan gum

## Inactive ingredients (Nighttime only)

anhydrous citric acid, FD&C blue #1, FD&C red #40, flavor, glycerin, polyethylene glycol, propylene glycol, purified water, sodium benzoate, sodium citrate dihydrate, sodium metabisulfite, sorbitol, sucralose, xanthan gum

### **Questions or comments?**

1-888-287-1915

### Principal display panel

#### equate™

Compare to Maximum Strength Mucinex® FAST-MAX® Day Time Severe Cold and Night Time Cold & Flu Active Ingredients<sup>†</sup>

NDC 49035-945-02

Maximum Strength <b>Daytime</b> <b>Severe Cold</b> <b>Acetaminophen</b> - Pain Reliever/Fever Reducer Dextromethorphan HBr - Cough Suppressant Guaifenesin - Expectorant Phenylephrine HCI - Nasal Decongestant <b>Multi-Symptom Relief</b> • Relieves aches, fever & sore throat • Controls cough • Relieves nasal & chest congestion • Thins & loosens mucus Ages 12+	Maximum Strength Nighttime Cold & Flu Acetaminophen - Pain Reliever/Fever Reducer Diphenhydramine HCI - Antihistamine/ Cough Suppressant Phenylephrine HCI - Nasal Decongestant Multi-Symptom Relief • Relieves aches, fever & sore throat • Controls cough • Relieves nasal congestion • Relieves runny nose & sneezing Ages 12+
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### TWO - 6 FL OZ (177 mL) BOTTLES

### TOTAL - 12 FL OZ (355 mL)

TAMPER EVIDENT: DO NOT USE IF IMPRINTED

### SAFETY SEAL UNDER CAP IS BROKEN OR MISSING

### **PARENTS:** Learn about teen medicine abuse: www.StopMedicineAbuse.org

## DISTRIBUTED BY: Walmart Inc., Bentonville, AR 72716

<sup>†</sup>This product is not manufactured or distributed by Reckitt Benckiser LLC, owner of the registered trademark Maximum Strength Mucinex ® FAST-MAX ® Day Time Severe Cold and Night Time Cold & Flu. REV0218C04201145 50844

W-2203-042011-45SS

Satisfaction guaranteed - Or we'll replace it or give you your money back. For guestions or comments or to report an undesired reaction or side effect, please call 1-888-287-1915.



Equate 44-042011

SEVERE COLD AND FLU acetaminophen, dextromethorphan hbr, diphenhydramine hcl, guaifenesin, phenylephrine hcl kit						
Product Information						
Р	Product Type         HUMAN OTC DRUG         Item Code (Source)         NDC:49035-945					
Packaging						
#	ltem Code	Package Description	Package Description		Marketing End Date	
1	NDC:49035-945- 02	in 1 PACKAGE; Type 0: Not a Combination roduct		08/15/2018		

Quantity Part #				tal Braduct Our stit	
			tal Product Quantity	У	
Part 1 1 BOTTLE					
Part 2 1 BOTTLE			177 mL		
Part 1 c	of 2				
CEV/EDE	COLD DAY				
			alfanaalia mbanudand		
acetaminop	onen, dextrometh	iorphan hbr, gu	aifenesin, phenyleph	nrine nci solution	
Product I	nformation				
Item Code		NDC:49035-842			
Route of A	dministration	ORAL			
Active Ing	gredient/Active	Moiety			
	Ingre	dient Name		Basis of Strength	n Strengtl
ACETAMINOPHEN (UNII: 36209ITL9D) (ACETAMINOPHEN - UNII: 36209ITL9D) ACETAMINOPHEN				ACETAMINOPHEN	650 mg in 20 mL
DEXTROMETHORPHAN HYDROBROMIDE (UNII: 9D2RTI9KYH)DEXTROMETHORPHAN(DEXTROMETHORPHAN - UNII:7355X3ROTS)HYDROBROMIDE					20 mg in 20 mL
GUAIFENESIN (UNII: 495W7451VQ) (GUAIFENESIN - UNII:495W7451VQ) GUAIFENESIN					400 mg in 20 mL
PHENYLEPHI UNII:1WS297V		IDE (UNII: 04JA59T	NSJ) (PHENYLEPHRINE -	PHENYLEPHRINE HYDROCHLORIDE	10 mg in 20 mL
	- ,				
Inactive I	ngredients				
	-	Ingredient	Name		Strength
ANHYDROUS	CITRIC ACID (UNII:	XF417D3PSL)			
EDETATE DISODIUM (UNII: 7FLD91C86K)					
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)					
FD&C RED NO. 40 (UNII: WZB9127XOA)					
GLYCERIN (UNII: PDC6A3C0OX)					
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)					
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)					
WATER (UNII: 059QF0KO0R)					
SODIUM BENZOATE (UNII: OJ245FE5EU)					
TRISODIUM CITRATE DIHYDRATE (UNII: B22547B95K)					
SODIUM METABISULFITE (UNII: 4VON5FNS3C)					
SORBITOL (UNII: 506T60A25R)					
SUCRALOSE					

<b>Product Charac</b>	teristics						
Color	blue	2	Score				
Shape			Size				
-		RY (Mixed)	Imprint	Code			
Contains							
Packaging							
# Item Code	Item Code Package Description			rketing Start Date		larketing End Date	
<b>1</b> NDC:49035-842- 45 Product 177 mL in 1 BOTTLE; Type 0: Not a Combination							
Marketing In	nformat	tion					
Marketing Category	Applica	ition Number or Monogra Citation	oh Ma	arketing Start Date	Marl	keting End Date	
OTC Monograph Drug	M012		08/1	5/2018			
		GHTTIME ramine hcl, phenylephrine	ncl soluti	on			
COLD AND F acetaminophen, d Product Inform	liphenhydr ation e)		ncl soluti	on			
COLD AND F acetaminophen, d Product Inform Item Code (Source Route of Administ	liphenhydr ation e) tration	NDC:49035-811 ORAL	ncl soluti	on			
COLD AND F acetaminophen, d Product Inform Item Code (Source Route of Administ	liphenhydr ation e) tration nt/Active	amine hcl, phenylephrine NDC:49035-811 ORAL Moiety	ncl soluti		nath	Strength	
COLD AND F acetaminophen, d Product Inform Item Code (Source Route of Administ Active Ingredie	liphenhydr ation e) tration nt/Active Ingre	NDC:49035-811 ORAL		on Basis of Stre ACETAMINOPHEN	ngth	650 mg	
COLD AND Fi acetaminophen, d Product Inform Item Code (Source Route of Administ Active Ingredien ACETAMINOPHEN (U DIPHENHYDRAMINE	liphenhydr ation e) tration nt/Active Ingre NII: 36209IT HYDROCHL	amine hcl, phenylephrine NDC:49035-811 ORAL Moiety dient Name L9D) (ACETAMINOPHEN - UNII:36 ORIDE (UNII: TC2D6JAD40)		Basis of Stre ACETAMINOPHEN DIPHENHYDRAMINE	-	650 mg in 20 mL 25 mg	
COLD AND F acetaminophen, d Product Inform Item Code (Source Route of Administ Active Ingredien ACETAMINOPHEN (U DIPHENHYDRAMINE (U PHENHYDRAMINE - U PHENYLEPHRINE HYD	liphenhydr ation e) tration nt/Active Ingre NII: 36209IT HYDROCHL JNII:8GTS825	amine hcl, phenylephrine NDC:49035-811 ORAL Moiety dient Name L9D) (ACETAMINOPHEN - UNII:36 ORIDE (UNII: TC2D6JAD40)	209ITL9D)	Basis of Stre ACETAMINOPHEN	-	in 20 mL	
Product Inform Item Code (Source Route of Administ Active Ingredien ACETAMINOPHEN (U DIPHENHYDRAMINE - U	liphenhydr ation e) tration nt/Active Ingre NII: 36209IT HYDROCHL JNII:8GTS825	amine hcl, phenylephrine NDC:49035-811 ORAL Moiety dient Name L9D) (ACETAMINOPHEN - UNII:36 ORIDE (UNII: TC2D6JAD40) 583M)	209ITL9D)	Basis of Stre ACETAMINOPHEN DIPHENHYDRAMINE HYDROCHLORIDE PHENYLEPHRINE	-	650 mg in 20 mL 25 mg in 20 mL 10 mg	
COLD AND Fi acetaminophen, d Product Inform Item Code (Source Route of Administ Active Ingredie ACETAMINOPHEN (U DIPHENHYDRAMINE - U PHENYLEPHRINE HY UNII: 1WS 297W6MV)	liphenhydr ation e) tration mt/Active Ingre NII: 36209IT HYDROCHL JNII:8GTS82: DROCHLOR	amine hcl, phenylephrine NDC:49035-811 ORAL Moiety dient Name L9D) (ACETAMINOPHEN - UNII:36 ORIDE (UNII: TC2D6JAD40) 583M) IDE (UNII: 04JA59TNSJ) (PHENYL	209ITL9D)	Basis of Stre ACETAMINOPHEN DIPHENHYDRAMINE HYDROCHLORIDE PHENYLEPHRINE		650 mg in 20 mL 25 mg in 20 mL 10 mg in 20 mL	
COLD AND Filacetaminophen, d Product Inform Item Code (Source Route of Administ Active Ingredien Active Ingredien (U) DIPHENHYDRAMINE (U) DIPHENHYDRAMINE (U) PHENYLEPHRINE HY UNII:1WS297W6MV) Inactive Ingredi	liphenhydr ation e) tration nt/Active Ingre NII: 36209IT HYDROCHL JNII:8GTS82: DROCHLOR	amine hcl, phenylephrine NDC:49035-811 ORAL Moiety dient Name L9D) (ACETAMINOPHEN - UNII:36 ORIDE (UNII: TC2D6JAD40) S83M) IDE (UNII: 04JA59TNSJ) (PHENYL Ingredient Name	209ITL9D)	Basis of Stre ACETAMINOPHEN DIPHENHYDRAMINE HYDROCHLORIDE PHENYLEPHRINE		650 mg in 20 mL 25 mg in 20 mL 10 mg	
COLD AND Fi acetaminophen, d Product Inform Item Code (Source Route of Administ Active Ingredie ACETAMINOPHEN (U DIPHENHYDRAMINE (U DIPHENHYDRAMINE - U PHENYLEPHRINE HY UNII: 1WS 297W6MV) Inactive Ingredi ANHYDROUS CITRIC	liphenhydr ation e) tration nt/Active Ingre NII: 36209IT HYDROCHL JNII:8GTS823 DROCHLOR ients ACID (UNII:	amine hcl, phenylephrine NDC:49035-811 ORAL Moiety dient Name L9D) (ACETAMINOPHEN - UNII:36 ORIDE (UNII: TC2D6JAD40) S83M) IDE (UNII: 04JA59TNSJ) (PHENYL Ingredient Name XF417D3PSL)	209ITL9D)	Basis of Stre ACETAMINOPHEN DIPHENHYDRAMINE HYDROCHLORIDE PHENYLEPHRINE		650 mg in 20 mL 25 mg in 20 mL 10 mg in 20 mL	
COLD AND Fi acetaminophen, d Product Inform Item Code (Source Route of Administ Active Ingredien ACETAMINOPHEN (U DIPHENHYDRAMINE (UPHENHYDRAMINE (U PHENYLEPHRINE HY UNII: 1WS 297W6MV) Inactive Ingredi ANHYDROUS CITRIC FD&C BLUE NO. 1 (U	ients ACID (UNII: JNII: H3R47K	amine hcl, phenylephrine NDC:49035-811 ORAL Moiety dient Name L9D) (ACETAMINOPHEN - UNII:36 ORIDE (UNII: TC2D6JAD40) 583M) IDE (UNII: 04JA59TNSJ) (PHENYL INGredient Name XF417D3PSL) 3TBD)	209ITL9D)	Basis of Stre ACETAMINOPHEN DIPHENHYDRAMINE HYDROCHLORIDE PHENYLEPHRINE		650 mg in 20 mL 25 mg in 20 mL 10 mg in 20 mL	
COLD AND F acetaminophen, d Product Inform Item Code (Source Route of Administ Active Ingredien ACETAMINOPHEN (U DIPHENHYDRAMINE (U PHENHYDRAMINE - U PHENYLEPHRINE HYD	ients ACID (UNII: JNII: H3R47K JNII: WZ B912	amine hcl, phenylephrine NDC:49035-811 ORAL Moiety dient Name L9D) (ACETAMINOPHEN - UNII:36 ORIDE (UNII: TC2D6JAD40) 583M) IDE (UNII: 04JA59TNSJ) (PHENYL INGredient Name XF417D3PSL) 3TBD)	209ITL9D)	Basis of Stre ACETAMINOPHEN DIPHENHYDRAMINE HYDROCHLORIDE PHENYLEPHRINE		650 mg in 20 mL 25 mg in 20 mL 10 mg in 20 mL	

		DC9Q167V3)				
VATER (UNII: 059QF0K00R)						
SODIUM BENZOATE (UNII: OJ245FE5EU)						
TRISODIUM CITRATE	DIHYD	RATE (UNII: B22547B95K)				
SODIUM METABISUL	.FITE (U	NII: 4VON5FNS3C)				
SORBITOL (UNII: 5067	T60A25R	)				
SUCRALOSE (UNII: 96	K6UQ3Z	D4)				
XANTHAN GUM (UNII: TTV12P4NEE)						
Product Charac	terist	ics				
Color	ore					
Shape			Siz	ze		
Flavor		BERRY (Mixed)	Im	print Code		
Contains						
Packaging						
# Item Code		Package Description		Marketing Start Date	Marketing Date	-
NDC:49035-811- 45     177 mL in 1 BOTTLE; Type 0: Not a Combination Product						
- 45 Pr	ouuci					
– 45 Pr	ouuci					
- 45 Pr	oduct					
Marketing In		nation				
- 45 Pr	oform	<b>nation</b> lication Number or Monograp Citation	h	Marketing Start Date	Marketin Date	-
Marketing In Category	oform	lication Number or Monograp	h			-
Marketing In Marketing	nform <sub>App</sub>	lication Number or Monograp	h	Date		-
Marketing In Category	nform <sub>App</sub>	lication Number or Monograp	h	Date		-
Marketing Marketing Category	App M012	lication Number or Monograp Citation	h	Date		-
Marketing In Marketing Category OTC Monograph Drug	M012	lication Number or Monograp Citation		Date		g End

# Labeler - Wal-Mart Stores Inc (051957769)

Establishment					
Name	Address	ID/FEI	<b>Business Operations</b>		
LNK International, Inc.		967626305	manufacture(49035-945) , pack(49035-945)		

Revised: 8/2023

Wal-Mart Stores Inc