WAL FLU SEVERE- acetaminophen, dextromethorphan hbr, phenylephrine hcl, diphenhydramine hcl Walgreen Company

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.

Walgreen Co. Wal-Flu[®] Severe Drug Facts

Active ingredients (in each packet) - Daytime

Acetaminophen 500 mg Dextromethorphan HBr 20 mg Phenylephrine HCl 10 mg

Purpose

Pain reliever/fever reducer Cough suppressant Nasal decongestant

Uses

- temporarily relieves these symptoms due to a cold:
- minor aches and pains
- minor sore throat pain
- headache
- nasal and sinus congestion
- cough due to minor throat and bronchial irritation
- temporarily reduces fever

Warnings

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

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

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.

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

- in a child under 12 years of age
- if you have ever had an allergic reaction to this product or any of its ingredients
- 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.

Ask a doctor before use if you have

- liver disease
- heart disease
- high blood pressure
- thyroid disease
- diabetes
- trouble urinating due to an enlarged prostate gland
- cough that occurs with too much phlegm (mucus)
- cough that lasts or is chronic such as occurs with smoking, asthma, or emphysema
- a sodium-restricted diet

Ask a doctor or pharmacist before use if you are

taking the blood thinning drug warfarin

When using this product

do not exceed recommended dosage

Stop use and ask a doctor if

- nervousness, dizziness, or sleeplessness occurs
- fever gets worse or lasts more than 3 days
- redness or swelling is present
- new symptoms occur
- pain, cough or nasal congestion gets worse or lasts more than 7 days
- 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: In case of overdose, get medical help or contact a Poison Control Center right away (1-800-222-1222). Prompt medical attention is critical for adults as well as for children even if you do not notice any signs or symptoms.

Directions

- do not use more than directed (see overdose warning)
- take every 4 hours, while symptoms persist. Do not take more than 6 packets in 24 hours unless directed by a doctor.

Age	Dose
adults and children 12 years of age and over	one packet
children under 12 years of age	do not use

- dissolve contents of one packet into 8 oz. hot water: sip while hot. Consume entire drink within 10-15 minutes.
- if using a microwave, add contents of one packet to 8 oz. of cool water: stir briskly before and after heating. Do not overheat.

Other information

- each packet contains: potassium 10 mg and sodium 25 mg
- **phenylketonurics:** contains phenylalanine 22 mg per packet
- store at 20-25°C (68-77°F). Protect product from heat and moisture.

Inactive ingredients

acesulfame potassium, anhydrous citric acid, aspartame, colloidal silicon dioxide, D&C yellow #10, FD&C blue #1, FD&C red #40, flavors, maltodextrin, pregelatinized starch, sodium citrate, sucrose, tribasic calcium phosphate

Questions or comments?

1-800-719-9260

Active ingredients (in each packet) - Nighttime

Acetaminophen 650 mg Diphenhydramine HCl 25 mg Phenylephrine HCl 10 mg

Purposes

Pain reliever/fever reducer Antihistamine/cough suppressant Nasal decongestant

Uses

- temporarily relieves these symptoms due to a cold:
- minor aches and pains
- minor sore throat pain
- headache

- nasal and sinus congestion
- runny nose
- sneezing
- itchy nose or throat
- itchy, watery eyes due to hay fever
- cough due to minor throat and bronchial irritation
- temporarily reduces fever

Warnings

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

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

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.

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

- in a child under 12 years of age
- if you have ever had an allergic reaction to this product or any of its ingredients
- with any other drug containing acetaminophen (prescription or nonprescription). If you are not sure whether a drug contains acetaminophen, ask a doctor or pharmacist.
- 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

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

• cough that lasts or is chronic such as occurs with smoking, asthma, or emphysema

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 exceed recommended dosage
- avoid alcoholic drinks
- marked drowsiness may occur
- 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 occurs
- fever gets worse or lasts more than 3 days
- redness or swelling is present
- new symptoms occur
- pain, cough or nasal congestion gets worse or lasts more than 7 days
- 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: In case of overdose, get medical help or contact a Poison Control Center right away (1-800-222-1222). Prompt medical attention is critical for adults as well as for children even if you do not notice any signs or symptoms.

Directions

- do not use more than directed (see overdose warning)
- take every 4 hours, while symptoms persist. Do not take more than 5 packets in 24 hours unless directed by a doctor.

Age	Dose
adults and children 12 years of age and over	one packet
children under 12 years of age	do not use

- dissolve contents of one packet into 8 oz. hot water: sip while hot. Consume entire drink within 10-15 minutes.
- if using a microwave, add contents of one packet to 8 oz. of cool water: stir briskly before and after heating. Do not overheat.

Other information

- each packet contains: potassium 10 mg and sodium 25 mg
- **phenylketonurics:** contains phenylalanine 13 mg per packet
- store at 20-25°C (68-77°F). Protect product from heat and moisture.

Inactive ingredients

acesulfame potassium, anhydrous citric acid, aspartame, colloidal silicon dioxide, D&C yellow #10, FD&C blue #1, FD&C red #40, flavors, maltodextrin, pregelatinized starch, sodium citrate, sucrose, tribasic calcium phosphate

Questions or comments?

1-800-719-9260

Package/Label Principal Display Panel

DAY & NIGHT PACK

MULTI-SYMPTOM

Wal-Flu[®]

Severe

COLD

ACETAMINOPHEN / PAIN RELIEVER / FEVER REDUCER

DEXTROMETHORPHAN HBr / COUGH SUPPRESSANT

PHENYLEPHRINE HCl / NASAL DECONGESTANT

Relieves nasal congestion, sore throat pain, cough, headache, body ache & fever

GREEN TEA & HONEY LEMON FLAVORS

6 PACKETS

Compare to Theraflu[®] Multi-Symptom Severe Cold & Theraflu[®] Nighttime Severe Cold & Cough active ingredients

NIGHTTIME

Wal-Flu[®]

Severe

COLD & COUGH

ACETAMINOPHEN / PAIN RELIEVER / FEVER REDUCER

DIPHENHYDRAMINE HCl / ANTIHISTAMINE / COUGH SUPPRESSANT

PHENYLEPHRINE HCl / NASAL DECONGESTANT

Relieves nasal congestion, cough, body ache, sore throat pain, runny nose, sneezing, headache & fever HONEY LEMON INFUSED WITH WHITE TEA FLAVORS

6 PACKETS

TOTAL 12 PACKETS



WAL FLU SEVERE

acetaminophen, dextromethorphan hbr, phenylephrine hcl, diphenhydramine hcl kit

PTU	duct Informat	tion			
Prod	uct T yp e	HUMAN OTC DRUG	Item Code ((Source)	NDC:0363-1319
Dack	aging				
#	Item Code	Package Descriptio	n	Marketing Start Date	Marketing End Date
1 ND	C:0363-1319-55	1 in 1 CARTON; Type 0: Not a Combin	nation Product	09/19/2018	
	ntity of Parts				
Qua	inity of I arts		Total Product Quantity		
-	5	Package Quantity		Total Product Qu	
Part	5	Package Quantity	6	Total Product Qu	
Part Part 1	#	Package Quantity	6 6	Total Product Qu	

Part 1 of 2

WAL FLU SEVERE COLD

acetaminophen, dextromethorphan hbr, phenylephrine hcl powder, for solution

Product Information	
Item Code (Source)	NDC:0363-8319
Route of Administration	ORAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ACETAMINOPHEN (UNII: 36209ITL9D) (ACETAMINOPHEN - UNII:36209ITL9D)	ACETAMINOPHEN	500 mg
DEXTRO METHO RPHAN HYDRO BRO MIDE (UNII: 9 D2RTI9 KYH) (DEXTRO METHO RPHAN - UNII: 7355X3ROTS)	DEXTROMETHORPHAN HYDROBROMIDE	20 mg
PHENYLEPHRINE HYDRO CHLO RIDE (UNII: 04JA59TNSJ) (PHENYLEPHRINE - UNII: 1WS297W6MV)	PHENYLEPHRINE HYDROCHLORIDE	10 mg

Inactive	Ingredients
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Ingredient Name	Strength
ACESULFAME POTASSIUM (UNII: 230V73Q5G9)	
ANHYDRO US CITRIC ACID (UNII: XF417D3PSL)	
ASPARTAME (UNII: Z0H242BBR1)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
D&C YELLOW NO. 10 (UNII: 35SW5USQ3G)	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	
FD&C RED NO.40 (UNII: WZB9127XOA)	
MALTODEXTRIN (UNII: 7CVR7L4A2D)	
SODIUM CITRATE, UNSPECIFIED FORM (UNII: 1Q73Q2JULR)	
SUCROSE (UNII: C151H8 M554)	
TRIBASIC CALCIUM PHOSPHATE (UNII: 91D9GV0Z28)	

Product Characteristics				
Color		Score		
Shape		Size		
Flavor	HONEY (green tea) , LEMON (green tea)	Imprint Code		
Contains				

Packaging			
# Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:0363-8319-91	6 in 1 CARTON; Type 0: Not a Combination Product		

	ormation				
Marketing Category		on Number or Monograph Citation	Marketing Start Da	ate Marketin	g End Date
OTC monograph final	part341		09/16/2018		
Part 2 of 2					
WAL FLU SE	VERE COI	LD AND COUGH			
		hydrochloride, phenylephrine hydro	ochloride powder fo	or solution	
uccuminoprien, up					
Product Informa	tion				
Item Code (Source) NDC:0363-0964					
Route of Administra	tion	ORAL			
Active Ingredien	t/Active Moi	ety			
	Ingr	edient Name	Basis o	f Strength	Strengtl
ACETAMINOPHEN (U	NII: 362O9ITL9I) (ACETAMINOPHEN - UNII:36209ITL9	D) ACETAMINOP	PHEN	650 mg
DIPHENHYDRAMINE UNII:8GTS82S83M)	HYDROCHLOR	IDE (UNII: TC2D6JAD40) (DIPHENHYDRA	AMINE - DIPHENHYDRA HYDROCHLOI		25 mg
PHENYLEPHRINE HY UNII:1WS297W6MV)	DRO CHLO RIDH	E (UNII: 04JA59TNSJ) (PHENYLEPHRINE	- PHENYLEPHRI HYDROCHLOI		10 mg
0111.1773237 W0147			IIIDROCIEOI		
Inactive Ingredie	nts				
		Ingredient Name		S	trength
ACESULFAME POTA	SSIUM (UNII: 23	OV73Q5G9)			
ANHYDRO US CITRIC	ACID (UNII: XF4	417D3PSL)			
ASPARTAME (UNII: ZO					
SILICON DIO XIDE (U		,			
	•	- ,			
D&C YELLOW NO. 1					
FD&C BLUE NO.1 (U					
FD&C BLUE NO. 1 (U FD&C RED NO. 40 (U	NII: WZB9127XC	A)			
FD&C BLUE NO. 1 (U FD&C RED NO. 40 (U MALTO DEXTRIN (UN	NII: WZB9127XC III: 7CVR7L4A2D	A))			
FD&C BLUE NO. 1 (U FD&C RED NO. 40 (U MALTO DEXTRIN (UN SODIUM CITRATE, U	NII: WZB9127XC III: 7CVR7L4A2D NSPECIFIED FC	A)			
FD&C BLUE NO. 1 (U FD&C RED NO. 40 (U MALTO DEXTRIN (UN SO DIUM CITRATE, U SUCROSE (UNII: C151)	NII: WZB9 127XC III: 7CVR7L4A2D NSPECIFIED FC H8 M554)	A)) RM (UNII: 1Q73Q2JULR)			
FD&C BLUE NO. 1 (U FD&C RED NO. 40 (U MALTO DEXTRIN (UN SODIUM CITRATE, U	NII: WZB9 127XC III: 7CVR7L4A2D NSPECIFIED FC H8 M554)	A)) RM (UNII: 1Q73Q2JULR)			
FD&C BLUE NO. 1 (U FD&C RED NO. 40 (U MALTO DEXTRIN (UN SODIUM CITRATE, U SUCROSE (UNII: C151) TRIBASIC CALCIUM	NII: WZB9127XC III: 7CVR7L4A2D NSPECIFIED FC H8M554) PHOSPHATE (U	A)) RM (UNII: 1Q73Q2JULR)			
FD&C BLUE NO. 1 (U FD&C RED NO. 40 (U MALTO DEXTRIN (UN SO DIUM CITRATE, U SUCROSE (UNII: C151) TRIBASIC CALCIUM	NII: WZB9 127XC III: 7CVR7L4A2D NSPECIFIED FC H8 M554) PHO SPHATE (U	A)) PRM (UNII: 1Q73Q2JULR) NII: 91D9GV0Z28)			
FD&C BLUE NO. 1 (U FD&C RED NO. 40 (U MALTO DEXTRIN (UN SO DIUM CITRATE, U SUCROSE (UNII: C151) TRIBASIC CALCIUM Product Characte Color WHITE	NII: WZB9 127XC III: 7CVR7L4A2D NSPECIFIED FC H8 M554) PHO SPHATE (U	A)) RM (UNII: 1Q73Q2JULR)	2	Score	
FD&C BLUE NO.1 (U) FD&C RED NO.40 (U) MALTODEXTRIN (UN SODIUM CITRATE, U) SUCROSE (UNI: C1512) TRIBASIC CALCIUM Product Character Color WHITE Shape (U)	NII: WZB9 127XC III: 7CVR7L4A2D NSPECIFIED FC H8 M554) PHO SPHATE (U	A)) PRM (UNII: 1Q73Q2JULR) NII: 91D9GV0Z28)	2	Size	
FD&C BLUE NO.1 (U) FD&C RED NO.40 (U) MALTODEXTRIN (UN) SODIUM CITRATE, U SUCROSE (UNI: C151) TRIBASIC CALCIUM Product Character Color WHITE Shape Flavor (U)	NII: WZB9 127XC III: 7CVR7L4A2D NSPECIFIED FC H8 M554) PHO SPHATE (U	A)) PRM (UNII: 1Q73Q2JULR) NII: 91D9GV0Z28)	2		de
FD&C BLUE NO.1 (U) FD&C RED NO.40 (U) MALTODEXTRIN (UN) SODIUM CITRATE, U SUCROSE (UNI: C1512) TRIBASIC CALCIUM Product Characte Color WHITE Shape	NII: WZB9 127XC III: 7CVR7L4A2D NSPECIFIED FC H8 M554) PHO SPHATE (U	A)) PRM (UNII: 1Q73Q2JULR) NII: 91D9GV0Z28)	2	Size	de
FD&C BLUE NO.1 (U) FD&C RED NO.40 (U) MALTODEXTRIN (UN) SODIUM CITRATE, U SUCROSE (UNI: C151) TRIBASIC CALCIUM Product Character Color WHITE Shape Flavor (U)	NII: WZB9 127XC III: 7CVR7L4A2D NSPECIFIED FC H8 M554) PHO SPHATE (U	A)) PRM (UNII: 1Q73Q2JULR) NII: 91D9GV0Z28)	2	Size	de

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0363-0964-91	6 in 1 CARTON; Type 0: Not a Combination Product		
M	arketing Info	rmation		
M	arketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ОТ	C monograph final	part341	11/05/2015	
M	arketing Info	rmation		
	arketing Info arketing Category	rmation Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
M	0		Marketing Start Date	Marketing End Date

Labeler - Walgreen Company (008965063)

Revised: 12/2019

Walgreen Company