ACETAMINOPHEN, CHLORPHENIRAMINE MALEATE, PHENYLEPHRINE HCLacetaminophen, chlorpheniramine maleate, phenylephrine hcl tablet OPMX LLC

Coldtac Ultra tabs

(Acetaminophen 500mg, Chlorpheniramine Maleate 4mg, Phenylephrine HCl 10mg Tablet)

Drug Facts

Active ingredients

Purpose

Active ingredients (in each caplet)	Purpose
Acetaminophen 500 mg	Pain reliever
Chlorpheniramine Maleate 4 mg	Antihistamine
Phenylephrine HCl 10 mg	Nasal decongestant

Uses

Temporarily relieves these symptoms of hay fever or other respiratory allergies:

- headache
- nasal congestion
- sinus congestion & pressure
- runny nose and sneezing
- minor aches & pain

Temporarily relieves these additional symptoms of hay fever:

- itching of the nose or throat
- itchy, watery eyes
- helps clear nasal passages
- helps decongest sinus opening and passages

Warnings

Liver warning:

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

- more than 6 tablets in 24 hours, which is the maximum daily amount for this product
- with other drugs contains 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 doctor promptly

Do not use

- with any other drug containing acetaminophen (prescription or nonprescription). If you are not sure whether a drug contains acetaminophen,
- 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
- if you have ever had an allergic reaction to this product or any of its ingredients.
- to make a child sleepy

Ask a doctor before use if you have:

- liver disease
- heart disease
- high blood pressure
- thyroid disease
- glaucoma
- diabetes
- trouble urinating due to enlarged prostate gland
- a breathing problem such as emphysema or chronic bronchitis

Ask a doctor ot pharmacist if you are:

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

When using this product:

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

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 last more than 3 days
- redness or swelling is present
- new symptoms occur

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. 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 take more than directed See overdose warning
- Swallow whole; do not crush, chew or dissolve
- Do not exceed 6 tablets per 24 hours

Adults and children 12 years of age &	1 tablet with water every 4-6 hours as
over	needed
Children 4 to 12 years of age	Ask a doctor
Children under 4 years of age	Do not use

Other information

- Store at room temperature between 20-25°C (68-77°F)
- Tamper evident: Do not use if pouch is torn, broken or shows any sing of tampering

Inactive ingredients

magnesium stearate, povidone, silicon dioxide, sodium benzoate, sodium starch glycolate, starch, talc

Questions or comments?

Call: (619) 600-5632 (Mon-Fri 9am-5pm EST) or https://www.opmx.us

Coldtac Ultra

NDC 69729-122-06

Rápido Alivio a las Molestias de la Gripe y Resfriado

- Dolor de Cabeza
- Estornudos
- Congestión Nasal

- Fiebre
- Escurrimiento Nasal Dolor de Garganta
- Cuerpo Cortado
- Ojos Llorosos

INSTRUCCIONES EN ESPAÑOL EN EL INTERIOR DE LA CAJA.

COLDTAC ULTRA

Tablets /Tabletas

COLDTAC

RXX

OLDTAC ULTR

NDC 69729-122-06

Acetaminophen 500 mg Chlorpheniramine Maleate 4 mg Phenylephrine HCl 10 mg

Effective and Quick Relief from Cold & Flu Symptoms

- Headache
- Fever
- Body Aches
- Sneezing
- * Runny Nose
- ♦ Watery Eyes♦ Nasal Congestion
- Sore Throat





Exclusively distributed by:



Chula Vista, CA 91910 Phone: 619-600-5632



Made for:



San Diego, CA

Questions or comments? Call: (619) 600-5632 (Mon-Fri 9am-5pm EST) or https://www.opmx.us

Inactive ingredients magnesium stearate, povidone, silicon dioxide, sodium benzoate, sodium benzoate,

Other information = Store at room temperature between 20°- 25°C (68°-77°F) = Avoid excessive heat and humidity = Tamper Evident: do not use if package or blister unit is open or shows any sign of tampering

	Children under 4 years of age
Ask a doctor	Children 4 to 12 years of age
4-6 hours as needed	
I tablet with water every	Adults and children

■ Swallow whole; do not črush, chew or dissolve ■ Do not exceed 6 tablets per 24 hours Do not take more than directed
 See Overdose warning

Directions

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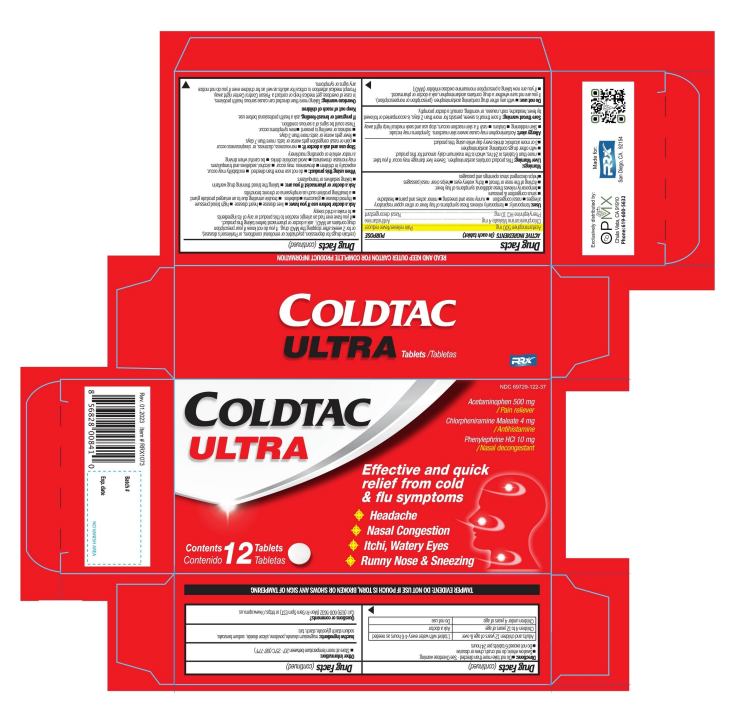


EXP. DATE

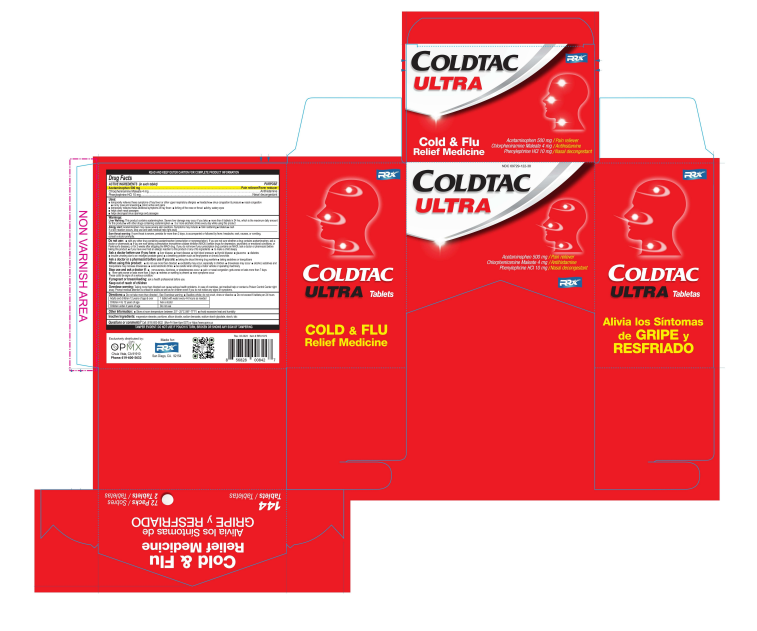
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Do not use: ■ with any other drug containing acetaminophen (prescription or nonprescription). If you are not sure monotamine oxidase inhibitor (MMO)) (eartain drugs for depression, psychiatric or emotional conditions, or Parkinson's disease), or for 2 weeks after stopping the MMOI drug. If you do not know if your prescription drug Parkinson's disease), or for 2 weeks after stopping the MMOI drug. If you do not know if your prescription drug
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Warnings Liver waming: Liver Warning: This product contains acetaminophen. Severe liver damage may occur if you take: ■ more than 6 tablets in 24 hrs., which is the maximum daily amount for this product may occur if you take: ■ more than 6 tablets in 24 hrs., which is the waximum daily amount for this product ■ with other drugs containing acetaminophen ■ 3 or more alcoholic drinks every day while using this product ■
Uses ■ temporarily relieves these symptoms of hay fever or other upper respiratory allergies ■ headache ■ sinus congestion & pressure ■ nasal congestion ■ runny nose and sneezing ■ minor aches and pains ■ temporarily relieves these additional symptoms of hay fever: ■ itching of the nose or throat ■ itchy, watery eyes ■ helps clear nasal passages ■ helps decongest sinus openings and passages ■ bloating ■ water-weight gain ■ headache ■ backache ■ muscle aches ■ fatigue
Active ingredients (in each caplet) Active ingredients (in each caplet) Actiminophen 500 mg. Chlorphenicamine Maleate 4 mg. Antihistamine Phenylephrine HCI 10 mg. Massil decongestant
Drug Facts

NDC 69729-122-37



NDC 69729-122-38



ACETAMINOPHEN, CHLORPHENIRAMINE MALEATE, PHENYLEPHRINE HCL

acetaminophen, chlorpheniramine maleate, phenylephrine hcl tablet

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:69729-122	
Route of Administration	ORAL			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
ACETAMINOPHEN (UNII: 36209ITL9D) (ACETAMINOPHEN - UNII: 36209ITL9D)	ACETAMINOPHEN	500 mg	
CHLORPHENIRAMINE MALEATE (UNII: V1Q0090J9Z) (CHLORPHENIRAMINE - UNII: 3U6I01965U)	CHLORPHENIRAMINE MALEATE	4 mg	
PHENYLEPHRINE HYDROCHLORIDE (UNII: 04JA59TNSJ) (PHENYLEPHRINE - UNII: 1W5297W6MV)	PHENYLEPHRINE HYDROCHLORIDE	10 mg	
UNII: 1W5 29 / WOMV)	HYDROCHLORIDE		

Inactive Ingredients			
Ingredient Name	Strength		
SODIUM BENZOATE (UNII: OJ245FE5EU)			
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)			
TALC (UNII: 7SEV7J4R1U)			
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)			
MAGNESIUM STEARATE (UNII: 70097M6I30)			
STARCH, CORN (UNII: O8232NY3SJ)			
POVIDONE K30 (UNII: U725QWY32X)			

Product Characteristics				
Color	white	Score	no score	
Shape	ROUND (BICONVEX)	Size	13mm	
Flavor		Imprint Code	S78	
Contains				

F	Packaging				
#	tem Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:69729-122- 27	2 in 1 POUCH; Type 0: Not a Combination Product	08/26/2019	10/24/2022	
2	NDC:69729-122- 37	12 in 1 POUCH; Type 0: Not a Combination Product	08/26/2019		
3	NDC:69729-122- 38	144 in 1 POUCH; Type 0: Not a Combination Product	08/26/2019		
4	NDC:69729-122- 06	3 in 1 CARTON	09/04/2023		
4		2 in 1 PACKET; 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	08/26/2019	
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Labeler - OPMX LLC (029918743)

Revised: 12/2024 OPMX LLC