# ZODRYL DEC 40 - codeine phosphate, guaifenesin and pseudoephedrine hydrochloride suspension CodaDose, Inc.

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.

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# **ZODRYL DEC 40 - codeine phosphate, guaifenesin and pseudoephedrine hydrochloride suspension**

#### **OTC - ACTIVE INGREDIENT**

Codeine phosphate 1 mg/1mL: cough suppressant; Guaifenesin 20 mg/1mL: expectorant; Pseudoephedrine hydrochloride 3.333 mg/1mL: decongestant

#### **PURPOSE**

Temporarily relieves: cough due to minor throat and bronchial irritation as may occur with the common cold or inhaled irritants; the intensity of coughing; the impulse to cough to help you go to sleep; temporarily relieves nasal congestion due to a cold; temporarily restores freer breathing through the nose; helps loosen phlegm (mucus) and thin bronchial passageways of bothersome mucus and makes coughs more productive

Warnings

#### **OTC - DO NOT USE**

in children who have chronic pulmonary disease, shortness of breath, or such as occurs with asthma or if cough is accompanied by excessive phlegm (mucus) unless directed by a doctor; if taking a monoamine oxidase inhibitor (MAOI)

#### OTC - ASK DOCTOR

if your child has glaucoma, a breathing problem such as emphysema or chronic bronchitis, heart disease, high blood pressure, thyroid disease, diabetes.

#### OTC - ASK DOCTOR/PHARMACIST SECTION

if you or your child are taking sedatives or tranquilizers; if you or your child are taking prescription MAOI (certain drugs for depression, psychiatric, or emotional conditions), or for 2 weeks after stopping the MAOI drug.

#### **OTC - WHEN USING THIS PRODUCT**

do not exceed recommended dosage; may cause or aggravate constipation; if nervousness, dizziness, or sleepiness occur, discontinue use and consult a doctor

#### OTC - STOP USE AND ASK A DOCTOR IF

cough persists for more than 1 week, tends to recur, or is accompanied by a fever, rash, or persistent headache. These could be signs of a serious condition.

#### OTC - KEEP THESE AND ALL DRUGS OUT OF REACH OF CHILDREN

In case of overdose, seek professional assistance for contact a Poison Control Center immediately.

#### Directions:

- Take every 4-6 hours
- Use only with enclosed calibrated oral dispenser
- Do not take more than 4 doses in 24 hours or as directed by a doctor

Children 2 to under 6 years of age: ask a doctor

Other information store at controlled room temperature 20°-25°C (68°-77°F).

#### **INACTIVE INGREDIENT**

Bittermask, citric acid, FD& C blue #1, FD& C red #40, galloquinate, glycerin, grape flavor, magnesium aluminometasilicate, methylparaben, purified water, sodium citrate dihydrate, sucralose, xanthan gum

#### **OTC – QUESTIONS SECTION**

Call 1-866-574-8861 24 hours a day, 7 days a week.

#### PACKAGE LABEL.PRINCIPAL DISPLAY PANEL

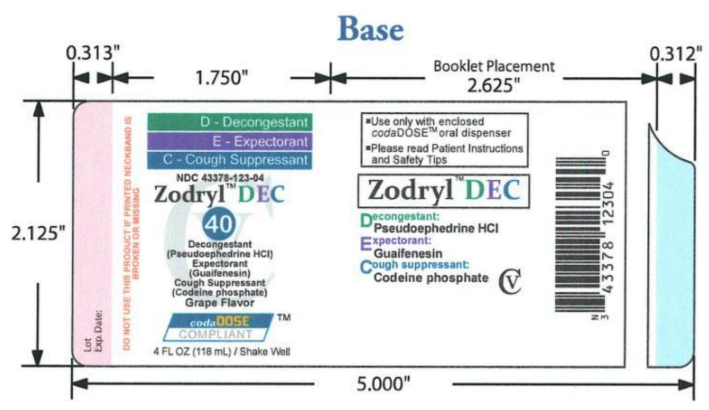


Figure 1. Primary Label- Front Page

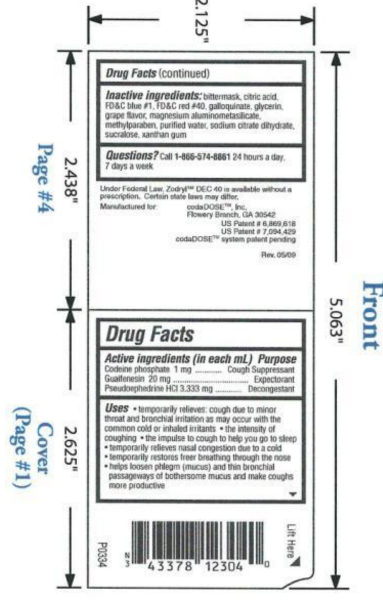


Figure 2. Primary Label - Second Page

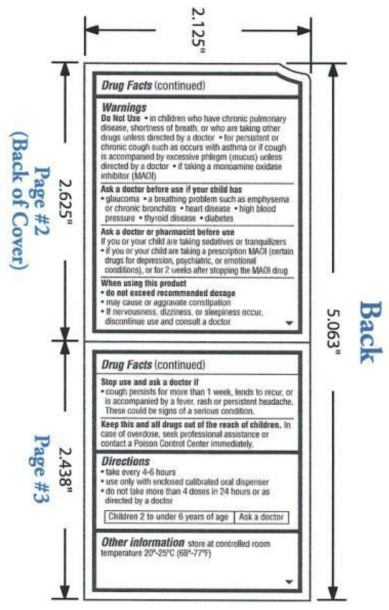


Figure 3. Primary Label – Last Page

#### **ZODRYL DEC 40**

codeine phosphate, guaifenesin and pseudoephedrine hydrochloride suspension

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:43378-123
Route of Administration	ORAL	DEA Sche dule	CV

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
PSEUDO EPHEDRINE HYDRO CHLO RIDE (UNII: 6 V9 V2RYJ8 N) (PSEUDO EPHEDRINE - UNII:7CUC9 DDI9 F)	PS EUDO EPHEDRINE HYDRO CHLO RIDE	14.998 mg in 4.5 mL	
CODEINE PHO SPHATE (UNII: GSL05Y1MN6) (CODEINE - UNII:Q830PW7520)	CODEINE PHOSPHATE	4.5 mg in 4.5 mL	
GUAIFENESIN (UNII: 495W7451VQ) (GUAIFENESIN - UNII:495W7451VQ)	GUAIFENESIN	90 mg in 4.5 mL	

Ingredient Name	Strength
ANHYDRO US CITRIC ACID (UNII: XF417D3PSL)	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	
FD&C RED NO. 40 (UNII: WZB9127XOA)	
TANNIC ACID (UNII: 28F9E0DJY6)	
GLYCERIN (UNII: PDC6A3C0OX)	
MAGNESIUM ALUMINUM SILICATE (UNII: 6M3P64V0NC)	
METHYLPARABEN (UNII: A2I8 C7HI9 T)	
WATER (UNII: 059QF0KO0R)	
SO DIUM CITRATE (UNII: 1Q73Q2JULR)	
SUCRALOSE (UNII: 96K6UQ3ZD4)	
XANTHAN GUM (UNII: TTV12P4NEE)	

Product Characteristics			
Color	purple	Score	
Shape		Size	
Flavor	GRAPE	Imprint Code	
Contains			

F	Packaging			
#	Item Code	Package Description	Marketing Start Date	<b>Marketing End Date</b>
1	NDC:43378-123-04	118 mL in 1 BOTTLE, PLASTIC		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part341	08/24/2009	

## Labeler - CodaDose, Inc. (831355115)

### **Registrant** - Gorbec Pharmaceutical Services Inc. (791919678)

Establishment				
Name	Address	ID/FEI	Business Operations	
Gorbec Pharmaceutical Services Inc.		79 19 19 6 78	manufacture	

Revised: 8/2009 CodaDose, Inc.