CHERRY THROAT DROPS- menthol lozenge Ricola USA 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.

CHERRY THROAT DROPS

Drug Facts

Purpose

Oral anesthetic

Active Ingredient (in each drop)

Menthol, 1.4 mg

Uses

temporarily relieves occasional minor irritation and pain associated with:

- sore mouth
- sore throat

Warnings

Do not use

• in children under 6 years of age unless directed by a doctor.

Stop use and ask a doctor if

- sore throat is severe, persists for more than 2 days, or is accompanied by fever, headache, rash, nausea or vomiting
- sore mouth symptoms do not improve in 7 days

Keep out of reach of children.

Directions

- adults and children 6 years and older: dissolve 2 drops (one at a time) slowly in the mouth. Do not bite or chew. Repeat every 2 hours as needed or as directed by a doctor
- children under 6 years: ask a doctor

Other Information

protect from heat and moisture

Inactive Ingredients

cherry concentrate, colors (carrot and hibiscus concentrate), extract of Ricola herb mixture (elder, horehound, hyssop, lemon balm, linden flowers, mallow, peppermint, sage, thyme, wild thyme), flavors (bitter alomnd and cherry), glucose syrup, malic acid, sugar



CHERRY THROAT DROPS menthol lozenge Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:63667-952 Route of Administration ORAL Active Ingredient/Active Moiety Ingredient Name Basis of Strength Strength

Inactive Ingredients			
Ingredient Name	Strength		
MALIC ACID (UNII: 817L1N4CKP)			
CHERRY (UNII: BUC5I9595W)			
SUCROSE (UNII: C151H8M554)			

Product Characteristics			
Color	red	Score	no score
Shape	OVAL	Size	24mm
Flavor	CHERRY (CHERRY)	Imprint Code	R
Contains			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:63667-952- 19	19 in 1 BAG; Type 0: Not a Combination Product	04/01/2023	
2	NDC:63667-952- 45	45 in 1 BAG; Type 0: Not a Combination Product	04/01/2023	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part356	04/01/2023	
	part356	04/01/2023	

Labeler - Ricola USA Inc. (177265261)

Establishment			
Name	Address	ID/FEI	Business Operations
Ricola Ag		485393768	manufacture(63667-952)

Revised: 4/2023 Ricola USA Inc.