DISCOUNT DRUG MART MAXIMUM STRENGTH URINARY PAIN RELIEFphenazopyridine hydrochloride tablet Reese Pharmaceutical Co

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

DRUG FACTS

Active ingredient (in each tablet)

Phenazopyridine Hydrochloride 97.5 mg .

Purpose

Urinary Analgesic

Warnings

Do not exceed recommended dosage

Ask doctor before use if you have

- kidney disease
- allergies to food, preservatives or dyes
- had a hypersensitive reaction to phenazopyridine

When using this product

■ stomach upset may occur, taking this product with or after meals may reduce stomach upset

■ your urine will become reddish-orange in color. This is not harmful, but care should be taken to avoid staining clothing or other items.

Stop use and ask doctor if

- your symptoms last for more than 2 days
- you suspect you are having an adverse reaction to the medication

If pregnant or breast feeding,

Ask a health professional before use.

Keep out of reach of children

In case of an overdose, get medical help or contact a Poison Control Center right away.

Use

Fast relief from urinary pain, burning, urgency and frequency associated with urinary tract infections.

Inactive ingredients

Lactose, magnesium silicate, magnesium stearate, microcrystalline cellulose, pharmaceutical glaze, and sodium starch glycolate.

Directions

- adults and children 12 years and over: take 2 tablets 3 times daily with a full glass of water, with or after meals as needed
- children under 12 years: consult a doctor
- Do not use for more than 2 days (12 tablets) without consulting a doctor



phenazopyridine hyd	lrochlori	ide table	et								
Product Informa	tion										
Product Type				HUMAN OTC DRUG			urce)	ז	NDC:10956-552		
Route of Administra	ntion		ORAL			ode (Source)			NDC.10330-332		
Route of Automistic	111011		ORIL								
Active Ingredien	t/Active	e Moie	ty								
	Ingre	Ingredient Name Basis						ength	Strength		
PHENAZO PYRIDINE HYDRO CHLO UNII:K2J09EMJ52)			IDE (UNII: 0 EWG668 W17) (PHENAZO PYRIDINE - PHENAZO PYRIDI HYDRO CHLO RID							97.5 mg	
Inactive Ingredie	ents										
Ingredient Name								Strength			
LACTOSE (UNII: J2B2A4N98G)											
MAGNESIUM SILICATE (UNII: 9B9691B2N9)											
Product Characteristics											
Color		red	Score			no			o score		
Shape		ROUND						7m			
Flavor			Imprint Code					j			
Contains											
Packaging											
# Item Code		Package Description			Marketing Start Date		te	Marketing End Date			
1 NDC:10956-552-12 1 in 1 CARTON				0 1/0 4			4/2010				
1 12 in 1 BLISTER PACK; Type 0: Not a Combination Product											
Marketing Information											
Marketing Category App			ation Number or Monograph Citation			Marketing Start Date			Marketing End Date		
unapproved drug other	pproved drug other		(0 1/0 4/20 10							

Labeler - Reese Pharmaceutical Co (004172052)

Registrant - Reese Pharmaceutical Co (004172052)

Establishment

Name	Address	ID/FEI	Business Operations
Reese Pharmaceutical Co		004172052	relabel(10956-552), repack(10956-552)