# QUALITY CHOICE URINARY PAIN RELIEF - phenazopyridine hydrochloride tablet Chain Drug Marketing Association

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.

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#### **DRUG FACTS**

#### **Active Ingredient**

Phenazopyridine Hydrochloride 95 mg

#### **Purpose**

**Urinary Analgesic** 

#### Uses

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

#### Warning

Do not exceed recommended dosage

## Ask Doctor before use if you have

- kidney disease
- allergies to foods, preservatives or dyes
- had a hypersensitive reaction to Phenazopyridine

# When using this product

- 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 a 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 the reach of children

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

#### **Directions**

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

## **Inactive Ingredients**

lactose, magnesium silicate, magnesium stearate, microcrystalline cellulose, pharmaceutical glaze, and sodium starch glycolate. May also contain: corn starch,

croscarmellose sodium, polyvinylpyrrolidone, pregelatinized starch and silicon dioxide.



## **QUALITY CHOICE URINARY PAIN RELIEF**

phenazopyridine hydrochloride tablet

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:83324-201
Route of Administration	ORAL		

Active Ingredient/Active Moiety		
Ingredient Name	<b>Basis of Strength</b>	Strength
PHENAZOPYRIDINE HYDROCHLORIDE (UNII: 0EWG668W17) (PHENAZOPYRIDINE - UNII: K2J09EMJ52)	PHENAZ OPYRIDINE HYDROCHLORIDE	95 mg

Inactive Ingredients		
Ingredient Name	Strength	
MAGNESIUM STEARATE (UNII: 70097M6I30)		
CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)		

SHELLAC (UNII: 46N107B710)

CROSCARMELLOSE SODIUM (UNII: M28OL1HH48)

SODIUM STARCH GLYCOLATE TYPE A CORN (UNII: AG9B65PV6B)

LACTOSE (UNII: J2B2A4N98G)

MAGNESIUM SILICATE (UNII: 9B9691B2N9)

<b>Product Characteris</b>	Product Characteristics			
Color	red	Score	no score	
Shape	ROUND	Size	7mm	
Flavor		Imprint Code	P95	
Contains				

Packaging			
# Item Code	Package Description	Marketing Start Date	Marketing End Date
NDC:83324- 201-30	1 in 1 CARTON	12/02/2025	
1	30 in 1 BLISTER PACK; Type 0: Not a Combination Product		

Marketing II	larketing Information		
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		12/02/2025	

# **Labeler -** Chain Drug Marketing Association (011920774)

Registrant - Reese Pharmaceutical Co (004172052)

Revised: 12/2025 Chain Drug Marketing Association