STOMACH RELIEF - MAXIMUM STRENGTH- bismuth subsalicylate liquid AptaPharma Inc.

Stomach Relief - Maximum Strength

Drug Facts

Active ingredient (in each 30 mL dose cup or 2 tablespoons)

Bismuth subsalicylate 1050 mg

Purposes

Bismuth subsalicylate Upset stomach reliever and antidiarrheal

Uses relieves ■ travelers' diarrhea ■ diarrhea ■ upset stomach due to overindulgence of food and drink including: ■ heart burn ■ indigestion ■ nausea ■ gas ■ belching

Warnings

Reye's syndrome: Children and teenagers who have or are recovering from chicken pox or flu-like symptoms should not use this product. When using this product, if changes in behavior with nausea and vomiting occur, consult a doctor because these symptoms could be an early sign of Reye's syndrome, a rare but serious illness.

Allergy alert: Contains salicylate. Do not take if you are

- allergic to salicylates (including aspirin)
- taking other salicylate products

Do not use if you have ■ an ulcer

■ a bleeding problem ■ black or bloody stool

Ask a doctor before use if you have

■ fever ■ mucus in stool

Ask a doctor or pharmacist before use if you are

taking any drug for ■ anticoagulation (thinning of the blood)

■ diabetes ■ gout ■ arthritis

When using this product a temporary, but harmless, darkening of the stool and/or tongue may occur

Slop use and ask a doctor if

- diarrhea lasts more than 2 days
- symptoms get worse or last more than 2 days
- ringing in the ears or loss of hearing occurs

If pregnant or breast feeding, ask health professional before use.

Keep out of reach of children.

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

Directions ■ Shake well before use

- use dose cup or tablespoon (TBSP)
- adults and children 12 years and over: 1 dose (30 mL or 2 TBSP) every 1 hour as needed
- do not exceed 4 doses (120 mL or 8 TBSP) in 24 hours
- use until diarrhea stops but not more than 2 days
- children under 12 years: ask a doctor
- drink plenty of clear fluids to help prevent dehydration caused by diarrhea

Other information

■ each 30 ml dose cup contains:

sodium 13 mg, salicylate 455 mg

■ protect from freezing ■ avoid excessive heat (over 104°F or 40°C) ■ low sodium

Inactive ingredients benzoic acid, D&C red #22, D&C red # 28, flavor, purified water, saccharin sodium, salicylic acid, sodium salicylate, xantham gum

Questions? 1-877-798-5944

Principal Display Panel

AP SAFE NDC 76281-538-26

*COMPARE TO the active ingredient in PEPTO-BISMOL™ MAXIMUM STRENGTH

Stomach Relief Bismuth Subsalicylate Antidiarrheal / Upset Stomach Reliever

Maximum Strength

- **5 Symptom Relief of:**
- Nausea Heartburn ●Indigestion
- Upset stomach Diarrhea

8 FL OZ (237 mL)

TAMPER EVIDENT: Do not use if imprinted shrinkband is missing or broken

*This product is not manufactured or distributed by Procter & Gamble, Inc., the distributor of Pepto-Bismol $^{\mathbb{M}}$.

Manufactured by: AptaPharma Inc.,

Made in USA

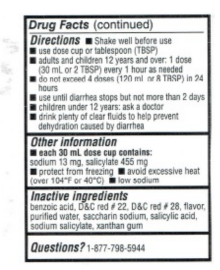
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STOMACH RELIEF - MAXIMUM STRENGTH

bismuth subsalicylate liquid

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:76281-538

Route of Administration ORAL

XANTHAN GUM (UNII: TTV12P4NEE)

Active Ingredient/Active Moiety

Ingredient Name

Basis of Strength

TE (UNII: 62TEY51RR1) (SALICYLIC ACID
BISMUTH

1050 mg

BISMUTH SUBSALICYLATE (UNII: 62TEY51RR1) (SALICYLIC ACID -

UNII:O414PZ4LPZ)

BISMUTH 1050 mg SUBSALICYLATE in 30 mL

Inactive Ingredients

Ingredient Name
Strength

BENZOIC ACID (UNII: 85KN0B0MIM)

D&C RED NO. 22 (UNII: 1678RKX8RT)

D&C RED NO. 28 (UNII: 767IP0Y5NH)

WATER (UNII: 059QF0K00R)

SACCHARIN SODIUM (UNII: SB8ZUX40TY)

SALICYLIC ACID (UNII: 0414PZ4LPZ)

SODIUM SALICYLATE (UNII: WQ1H85SYP)

Product Characteristics			
Color	pink (color suspension)	Score	
Shape		Size	
Flavor	MINT	Imprint Code	
Contains			

Packaging				
# Ite	m Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:	76281-538-	237 mL in 1 BOTTLE; Type 0: Not a Combination Product	10/01/2018	

Marketing Information				
Marketing Application Number or Monograph Category Citation		Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M008	10/01/2018		

Labeler - AptaPharma Inc. (790523323)

Registrant - AptaPharma Inc. (790523323)

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Name Address ID/FEI Business Operations

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AptaPharma Inc.	-	790523323	manufacture(76281-538)

Revised: 12/2023 AptaPharma Inc.